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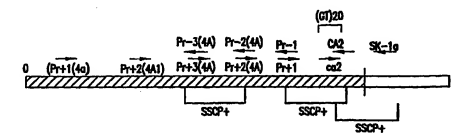
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(57) Abstract

In a preferred aspect of the invention, the upstream sequences of the TIGR protein encoding sequence can be used to diagnose a sensitivity to steroids and a risk for glaucoma or ocular hypertensive disorders. Methods, kits, and nucleic acids containing polymorphisms, base substitutions, or base additions located within the upstream region and within protein—encoding regions of the TIGR gene are also provided. The upstream sequences disclosed, including the TIGR promoter regions and those regions possessing functional characteristics associated with or possessed by the TIGR gene 5' regulatory region, can also be used to generate cells, vectors, transgenic animals, and nucleic acid constructs useful in a variety of diagnostic and prognostic methods and kits as well as therapeutic compounds, compositions and methods.

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NUCLEIC ACIDS, KITS, AND METHODS FOR THE DIAGNOSIS, PROGNOSIS AND TREATMENT OF GLAUCOMA AND RELATED DISORDERS

FIELD OF THE INVENTION

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The present invention relates to the field of diagnostic and prognostic methods and kits, treatments, and compositions useful in understanding and identifying glaucoma, related intraocular pressure-disorders, and steroid sensitivity.

CROSS REFERENCE TO RELATED APPLICATIONS

This application is a continuation-in-part of U.S. Patent Application serial no. 09/227,881, filed January 11, 1999, specifically incorporated by reference herein, which is a continuation-in-part of U.S. Patent Application serial no. 08/938,669, filed September 26, 1997, specifically incorporated by reference herein, which is a continuation-in-part of U.S. Patent Application serial no. 08/791,154, filed January 28, 1997, also specifically incorporated by reference herein.

BACKGROUND OF THE INVENTION

A group of debilitating eye diseases, the "Glaucomas" represent the leading cause of preventable blindness in the United States and other developed nations. In general, glaucomas are characterized by the alteration of the trabecular meshwork (TM), which consists of specialized endothelial cells and their associated connective tissue. The TM endothelial cells line the path the aqueous humor of the eye filters through during the normal, physiological flux. The cells generate and regulate the TM by producing extracellular molecules, the composition of which is thought to directly control the aqueous fluid flow.

In Primary Open Angle Glaucoma ("POAG"), the most common form of glaucoma, an alteration in the TM leads to an obstruction of the normal ability of aqueous humor to leave its chamber surrounding the iris. However, the specific cells in the chamber between the iris and the comea, in a region called the iridocomeal angle, remain "open" in that they continue to allow the egress of aqueous fluid (see, Vaughan, D. et al., In: General Ophthalmology, Appleton & Lange, Norwalk, CT, pp. 213-230 (1992); and Gray's Anatomy, 37th Ed., Churchill Livingstone, London, pp. 1180-1190 (1989)). As a result of the alteration in the TM and the obstruction, an increased intraocular pressure ("IOP") can be observed. IOP can result in progressive visual loss and blindness if not treated appropriately and in a timely fashion.

Glaucomas are estimated to affect between 0.4% and 3.3% of all adults over 40 years old (Leske, M.C. et al., Amer. J. Epidemiol. 113:1843-1846 (1986); Bengtsson, B., Br. J. Ophthamol. 73:483-487 (1989); Strong, N.P., Ophthal. Physiol. Opt. 12:3-7 (1992)). Moreover, the prevalence of the disease rises to over 6% of those 75 years or older (Strong, N.P., Ophthal. Physiol. Opt. 12:3-7 (1992)).

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A link between steroid, corticosteroid, or glucocorticoid treatments and the increased IOP found in POAG disease has long been suspected. While only 5% of the normal population have high IOP increases in response to topical glucocorticoids, greater than 40-50% of similarly treated patients with POAG show a high IOP increase (16 mm Hg). In addition, an Open Angle Glaucoma may be induced by exposure to glucocorticoids. This observation has suggested that an increased or abnormal glucocorticoid response in trabecular cells of the TM may be involved in POAG (Zhan, G.L. et al., Exper. Eye Res. 54:211-218 (1992); Yun, A.J. et al., Invest. Ophthamol. Vis. Sci. 30:2012-2022 (1989); Clark, A.F., Exper. Eye Res. 55:265 (1992); Klemetti, A., Acta Ophthamol. 68:29-33 (1990); Knepper, P.A., U.S. Patent No. 4,617,299).

The ability of glucocorticoids to induce a glaucoma-like condition has led to efforts to identify genes or gene products induced by the cells of the trabecular meshwork in response (Polansky, J.R. et al., In: Glaucoma Update IV, Springer-Verlag, Berlin, pp. 20-29 (1991); Polansky J.R. and Weinrob, R.N., In: Handbook of Experimetal Pharmacology, Vol. 69, Springer-Verlag, Berlin, pp. 461-538 (1984)). Initial efforts using short-term exposure to dexamethasone revealed only changes in specific protein synthesis. Extended exposure to relatively high levels of dexamethasone was, however, found to induce the expression of related 66 kD and 55 kD proteins that could be visualized by gel electrophoresis (Polansky, J.R. et al., In: Glaucoma Update IV, Springer-Verlag, Berlin, pp. 20-29 (1991)). The induction kinetics of these proteins as well as their dose response characteristics were similar to the kinetics that were required for steroid-induced IOP elevation in human subjects (Polansky, J.R. et al., In: Glaucoma Update IV, Springer-Verlag, Berlin, pp. 20-29 (1991)). Problems of aggregation and apparent instability or loss of protein in the purification process were obstacles in obtaining a direct protein sequence.

Nguyen et al., U.S. Patent Application No: 08/649,432, filed May 17, 1996, now U.S. Patent No. 5,789,169, the entire disclosure of which is hereby incorporated by reference as if set forth at length herein, disclosed a novel protein sequence (the TIGR, trabecular meshwork inducible glucocorticoid response protein) highly induced by glucocorticoids in the endothelial lining cells of the human trabecular meshwork. Nguyen et al. also disclosed the cDNA sequence for that protein, the protein itself, molecules that bind to it, and nucleic acid molecules that

encode it, and provided improved methods and reagents for diagnosing glaucoma and related disorders, as well as for diagnosing other diseases or conditions, such as cardiovascular, immunological, or other diseases or conditions that affect the expression or activity of the protein.

Because increased IOP is a readily measurable characteristic of glaucoma, the diagnosis of the disease is largely screened for by measuring intraocular pressure (tonometry) (Strong, N.P., Ophthal. Physiol. Opt. 12:3-7 (1992), Greve, M. et al., Can. J. Ophthamol. 28:201-206 (1993)). Unfortunately, because glaucomatous and normal pressure ranges overlap, such methods are of limited value unless multiple readings are obtained (Hitchings, R.A., Br. J. Ophthamol. 77:326 (1993); Tuck, M.W. et al., Ophthal. Physiol. Opt. 13:227-232 (1993); Vaughan, D. et al., In: General Ophthamology, Appleton & Lange, Norwalk, CT, pp. 213-230 (1992); Vernon, S.A., Eye 7:134-137 (1993)). Patients may also have a differential sensitivity to optic nerve damage at a given IOP. For these reasons, additional methods, such as direct examination of the optic disk and determination of the extent of a patient's visual field loss are often conducted to improve the accuracy of diagnosis (Greve, M. et al., Can. J. Ophthamol. 28:201-206 (1993)). Moreover, these techniques are of limited prognostic value. In some aspects, the present invention fulfills the need for improved diagnostic and prognostic methods.

The elevation of intraocular pressure (IOP) due to topical corticosteroids (and other routes of administration) is an important clinical problem that limits the clinical use of these effective anti-inflammatory agents. If not observed in sufficient time, the IOP elevation (especially in certain individuals who show the high end of steroid-induced IOP elevations) can result in optic nerve damage and permanent visual field loss, termed "steroid glaucoma." Even patients taking inhaled, nasal, rectal, and facial steroids may be at risk. The present invention, in part, provides improved diagnostic agents, prognostic agents, therapeutic agents and methods that address this clinical problem.

SUMMARY OF THE INVENTION

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The invention relates to nucleic acids, genes, proteins and cells that can be used in the treatment, diagnosis, prognosis, and identification of glaucoma, IOP-related disorders, or steroid sensitivity. The invention encompasses numerous agents, compositions, and methods, some of which are described by the objects and aspects of the invention detailed below. Others can be devised from the entire contents of this disclosure, and from the detailed description that follows.

In one aspect, the invention relates to nucleic acids comprising non-coding regions or promoter regions associated with the TIGR (trabecular meshwork inducible glucocorticoid

response) gene of mammals. These nucleic acids can be used in identifying polymorphisms in the genomes of mammals and humans that predict steroid sensitivity or a susceptibility to glaucomas or diseases related to alterations in IOP. A number of diagnostic or prognostic methods and kits can be designed from these nucleic acids.

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In one embodiment, the nucleic acids can be used to identify or detect a single base polymorphism in a genome. In other embodiments, two or more single base polymorphisms or multiple base polymorphisms can be identified or detected. The detection of a known polymorphism can be the basis for diagnostic and prognostic methods and kits of the invention. Various methods of detecting nucleic acids can be used in these methods and with the kits, including, but not limited to, solution hybridization, hybridization to microarrays containing immobilized nucleic acids or other immobilized nucleic acids, amplification-based methods such as PCR and the like, and an appropriate biosensor apparatus comprising a nucleic acid or nucleic acid binding reagent.

In another aspect, the invention relates to specific sequences and variants or mutants from the promoter or 5' regulatory region of the human TIGR gene and nucleic acids incorporating these sequences, variants or mutants. The nucleic acids can be incorporated into the methods and kits of the invention, or used in expression systems, vectors, and cells to produce a protein or polypeptide of interest, or used in methods to identify or detect regulatory proteins or proteins that specifically bind to promoter or regulatory regions of the TIGR gene. While many of the examples below detail work from human tissue, other animals may be used as a source of the sequences.

In one embodiment of this aspect of the invention, for example, nucleic acids having the disclosed TIGRmt11 sequence variant, represented by the change at nucleotide 5113 in SEQ ID NO: 1, 3, or 34 from T to C, or the change in nucleotide 5117 in SEQ ID NO: 2 from T to C. The presence of sequence variant mt11 is linked to the high IOP response to steroid treatments and a nucleic acid incorporating the singe base substitution can be used to identify and determine individuals at risk for developing glaucoma from undergoing a steroid treatment therapy, or a progression from an ocular hypertensive state, or those with a steroid sensitivity. And, because of the link between high IOP responses to steroids and the later development of glaucoma, nucleic acids having the TIGRmt11 sequence variant may also be used to identify the risk of developing glaucomas, such as POAG. The identification of changes in IOP can be done by any known means, however, the "Armaly" criteria is preferred (see Armaly, M.F., Arch. Ophthalmol. 70:492 (1963); Armaly, M.F., Arch Ophtalmol. 75:32-35 (1966); Kitazawa, Y. et al., Arch. Ophthalmol. 99:819-823 (1981); Lewis, J.M. et al., Amer. J. Ophthalmol. 106:607-612 (1988);

Becker, B. et al. Amer. J.Ophthalmol. 57:543 (1967), all of which are specifically incorporated herein by reference in their entireties).

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An object of the invention is to provide a method for diagnosing glaucoma in a patient which comprises the steps: (A) incubating under conditions permitting nucleic acid hybridization: a marker nucleic acid molecule, said marker nucleic acid molecule comprising a nucleotide sequence of a polynucleotide that specifically hybridizes to a polynucleotide that is linked to a TIGR promoter, and a complementary nucleic acid molecule obtained from a cell or a bodily fluid of said patient, wherein nucleic acid hybridization between said marker nucleic acid molecule, and said complementary nucleic acid molecule obtained from said patient permits the detection of a polymorphism whose presence is predictive of a mutation affecting TIGR response in said patient; (B) permitting hybridization between said marker nucleic acid molecule and said complementary nucleic acid molecule obtained from said patient; and (C) detecting the presence of said polymorphism, wherein the detection of the polymorphism is diagnostic of glaucoma.

Another object of the invention is to provide a method for prognosing glaucoma in a patient which comprises the steps: (A) incubating under conditions permitting nucleic acid hybridization: a marker nucleic acid molecule, said marker nucleic acid molecule comprising a nucleotide sequence of a polynucleotide that specifically hybridizes to a polynucleotide that is linked to a TIGR promoter, and a complementary nucleic acid molecule obtained from a cell or a bodily fluid of said patient, wherein nucleic acid hybridization between said marker nucleic acid molecule, and said complementary nucleic acid molecule obtained from said patient permits the detection of a polymorphism whose presence is predictive of a mutation affecting TIGR response in said patient; (B) permitting hybridization between said marker nucleic acid molecule and said complementary nucleic acid molecule obtained from said patient; and (C) detecting the presence of said polymorphism, wherein the detection of the polymorphism is prognostic of glaucoma.

Another object of the invention is to provide marker nucleic acid molecules capable of specifically detecting TIGRmt1, TIGRmt2, TIGRmt3, TIGRmt4, TIGRmt5, TIGRmt11 and TIGRsv1.

Another object of the invention is to provide a method for diagnosing steroid sensitivity in a patient which comprises the steps: (A) incubating under conditions permitting nucleic acid hybridization: a marker nucleic acid molecule, the marker nucleic acid molecule comprising a nucleotide sequence of a polynucleotide that is linked to a TIGR promoter, and a complementary nucleic acid molecule obtained from a cell or a bodily fluid of the patient, wherein nucleic acid hybridization between the marker nucleic acid molecule, and the complementary nucleic acid molecule obtained from the patient permits the detection of a polymorphism whose presence is predictive of a mutation affecting TIGR response in the patient; (B) permitting hybridization between said TIGR-encoding marker nucleic acid molecule and the complementary nucleic acid

molecule obtained from the patient; and (C) detecting the presence of the polymorphism, wherein the detection of the polymorphism is diagnostic of steroid sensitivity.

Further objects of the invention provide a nucleic acid molecule that comprises the sequence of SEQ ID NO: 1 or 34, recombinant DNA molecules containing a polynucleotide that specifically hybridizes to SEQ ID NO: 1 or 34 and substantially purified molecules that specifically bind to a nucleic acid molecule that comprises the sequence of SEQ ID NO: 1 or 34.

Further objects of the invention provide a nucleic acid molecule that comprises the sequence of SEQ ID NO: 3, recombinant DNA molecules containing a polynucleotide that specifically hybridizes to SEQ ID NO: 3 and substantially purified molecules that specifically bind to a nucleic acid molecule that comprises the sequence of SEQ ID NO: 3.

Additional objects of the invention provide a nucleic acid molecule that comprises the sequence of SEQ ID NO: 4, recombinant DNA molecules containing a polynucleotide that specifically hybridizes to SEQ ID NO: 4 and substantially purified molecules that specifically bind to a nucleic acid molecule that comprises the sequence of SEQ ID NO: 4.

Additional objects of the invention provide a nucleic acid molecule that comprises the sequence of SEQ ID NO: 5, recombinant DNA molecules containing a polynucleotide that specifically hybridizes to SEQ ID NO: 5 and substantially purified molecules that specifically bind to a nucleic acid molecule that comprises the sequence of SEQ ID NO: 5.

An additional object of the present invention is to provide a method of treating glaucoma which comprises administering to a glaucomatous patient an effective amount of an agent that inhibits the synthesis of a TIGR protein.

Indeed, the molecules of the present invention may be used to diagnose diseases or conditions which are characterized by alterations in the expression of extracellular proteins.

BRIEF DESCRIPTION OF THE FIGURES:

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Figures 1A, 1B, 1C, 1D and 1E provide the nucleic acid sequence of a TIGR promoter region (SEQ ID NO: 1) from an individual without glaucoma.

Figures 2A, 2B, 2C and 2D provide the location and sequence changes highlighted in bold associated with glaucoma mutants TIGRmt1, TIGRmt2, TIGRmt3, TIGRmt4, TIGRmt5, TIGRmt11, and TIGRsv1 (SEQ ID NO: 2).

Figures 3A, 3B, 3C, 3D, 3E, 3F, and 3G provide nucleic acid sequences of a TIGR promoter, and TIGR exons, TIGR introns and TIGR downstream sequences (SEQ ID NO: 3, SEQ ID NO: 4, and SEQ ID NO: 5).

Figure 4 provides a diagrammatic representation of the location of primers on the TIGR gene promoter for Single Strand Conformational Polymorphism (SSCP) analysis.

Figure 5 provides a diagrammatic representation of the TIGR exons and the arrangement of SSCP primers.

Figure 6 provides a homology analysis of TIGR homology with olfactomedin and olfactomedin-related proteins.

Figure 7 shows the nucleotide sequence of TIGR (SEQ ID NO: 26).

Figure 8 shows the amino acid sequence of TIGR (SEQ ID NO: 32).

Figure 9 shows the results of a gel shift assay and a diagramatic representation of the results. Lanes labeled "TM" represent the binding of cellular components from human trabecular meshwork cells to the 283 base pair TIGR 5' regulatory region. Lanes labeled "HeLa" represent the binding of cellular components from HeLa cells to the same nucleic acid. A much higher amount of cellular component binds in the +DEX (dexamethasone) treated TM cells than in the -DEX Tm cells (no dexamethsone).

DETAILED DESCRIPTION OF THE INVENTION

I. Agents of the Invention

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As used herein, the term "glaucoma" has its art recognized meaning, and includes both primary glaucomas, secondary glaucomas, juvenile glaucomas, congenital glaucomas, and familial glaucomas, including, without limitation, pigmentary glaucoma, high tension glaucoma and low tension glaucoma and their related diseases. The methods of the present invention are particularly relevant to the diagnosis of POAG, OAG, juvenile glaucoma, and inherited glaucomas. The methods of the present invention are also particularly relevant to the prognosis of POAG, OAG, juvenile glaucoma, and inherited glaucomas. A disease or condition is said to be related to glaucoma if it possesses or exhibits a symptom of glaucoma, for example, an increased intra-ocular pressure resulting from aqueous outflow resistance (see, Vaughan, D. et al., In: General Ophthamology, Appleton & Lange, Norwalk, CT, pp. 213-230 (1992)). The preferred agents of the present invention are discussed in detail below.

The agents of the present invention are capable of being used to diagnose the presence or severity of glaucoma and its related diseases in a patient suffering from glaucoma (a "glaucomatous patient"). The agents of the present invention are also capable of being used to prognose the presence or severity of glaucoma and its related diseases in a person not yet suffering from any clinical manifestations of glaucoma. Such agents may be either naturally occurring or non-naturally occurring. As used herein, a naturally occurring molecule may be "substantially purified," if desired, such that one or more molecules that is or may be present in a naturally occurring preparation containing that molecule will have been removed or will be present at a lower concentration than that at which it would normally be found.

The agents of the present invention will preferably be "biologically active" with respect to either a structural attribute, such as the capacity of a nucleic acid to hybridize to another nucleic acid molecule, or the ability of a protein to be bound by antibody (or to compete with another molecule for such binding). Alternatively, such an attribute may be catalytic, and thus involve the capacity of the agent to mediate a chemical reaction or response.

As used herein, the term "TIGR protein" refers to a protein having the amino acid sequence of SEQ ID NO: 32. As used herein, the agents of the present invention comprise nucleic acid molecules, proteins, and organic molecules.

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As indicated above, the trabecular meshwork has been proposed to play an important role in the normal flow of the aqueous, and has been presumed to be the major site of outflow resistance in glaucomatous eyes. Human trabecular meshwork (HTM) cells are endothelial like cells which line the outflow channels by which aqueous humor exits the eye; altered synthetic function of the cells may be involved in the pathogenesis of steroid glaucoma and other types of glaucoma. Sustained steroid treatment of these cells are interesting because it showed that a major difference was observed when compared to 1-2 day glucocorticoid (GC) exposure. This difference appears relevant to the clinical onset of steroid glaucoma (1-6 weeks).

Although trabecular meshwork cells had been found to induce specific proteins in response to glucocorticoids (see, Polansky, J.R., In: "Basic Aspects of Glaucoma Research III", Schattauer, New York 307-318 (1993)), efforts to purify the expressed protein were encumbered by insolubility and other problems. Nguyen, T.D. et al., (In: "Basic Aspects of Glaucoma Research III", Schattauer, New York, 331-343 (1993), herein incorporated by reference) used a molecular cloning approach to isolate a highly induced mRNA species from glucocorticoid-induced human trabecular cells. The mRNA exhibited a time course of induction that was similar to the glucocorticoid-induced proteins. The clone was designated "II.2" (ATCC No: 97994, American Type Culture Collection, Manassas, VA).

Nguyen et al., U.S. Patent Application No: 08/649,432 filed May 17, 1996, isolated a II.2 clone which encoded a novel secretory protein that is induced in cells of the trabecular meshwork upon exposure to glucocorticoids. It has been proposed that this protein may become deposited in the extracellular spaces of the trabecular meshwork and bind to the surface of the endothelial cells that line the trabecular meshwork, thus causing a decrease in aqueous flow. Quantitative dot blot analysis and PCR evaluations have shown that the mRNA exhibits a progressive induction with time whereas other known GC-inductions from other systems and found in HTM cells (metallothionein, alpha-1 acid glycoprotein and alpha-1 antichymotrypsin) reached maximum level at one day or earlier. Of particular interest, the induction level of this clone was very high (4-6% total cellular mRNA) with control levels undetectable without PCR method. Based on studies of 35S methionine cell labeling, the clone has the characteristics recently

discovered for the major GC-induced extracellular glycoprotein in these cells, which is a sialenated, N-glycosylated molecule with a putative inositol phosphate anchor. The induction of mRNA approached 4% of the total cellular mRNA. The mRNA increased progressively over 10 days of dexamethasone treatment. The II.2 clone is 2.0 Kb whereas the Northern blotting shows a band of 2.5 Kb. Although not including a poly A tail, the 3' end of the clone contains two consensus polyadenylation signals.

A genomic clone was isolated and designated P₁TIGR clone (ATCC No: 97570, American Type Culture Collection, Rockville, Maryland). In-situ hybridization using the P₁TIGR clone shows a TIGR gene and/or a sequence or sequences that specifically hybridize to the TIGR gene located at chromosome 1, q21-27, and more preferably to the TIGR gene located at chromosome 1, q22-26, and most preferably to the TIGR gene located at chromosome 1, q24. Clone P₁TIGR comprises human genomic sequences that specifically hybridize to the TIGR gene cloned into the *Bam*HI site of vector pCYPAC (Ioannou *et al.*, *Nature Genetics*, 6:84-89 (1994) herein incorporated by reference).

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As used herein, the term "TIGR gene" refers to the region of DNA involved in producing a TIGR protein; it includes, without limitation, regions preceeding and following the coding region as well as intervening sequences between individual coding regions.

As used herein, the term "TIGR exon" refers to any interrupted region of the TIGR gene that serves as a template for a mature TIGR mRNA molecule. As used herein, the term "TIGR intron" refers to a region of the TIGR gene which is non-coding and serves as a template for a TIGR mRNA molecule.

Localization studies using a Stanford G3 radiation hybrid panel mapped the TIGR gene near the D1S2536 marker with a LOD score of 6.0 (Richard et al., American Journal of Human Genetics 52.5: 915-921 (1993), herein incorporated by reference); Frazer et al., Genomics 14.3: 574-578 (1992), herein incorporated by reference; Research Genetics, Huntsville, Alabama). Other markers in this region include: D1S210; D1S1552; D1S2536; D1S2790; SHGC-12820; and D1S2558.

Sequences located upstream of the TIGR coding region are isolated and sequenced in a non-glaucomic individual. The upstream sequence is set forth in SEQ ID. No. 1 and 34. Sequence comparisons of the upstream region of a non-glaucoma individual and individuals with glaucoma identify a number of mutations in individuals with glaucoma. Some of these mutations are illustrated in Figure 2, the sequence of which can be used to identify the exact changes in the human genomic sequences from the upstream region of the TIGR gene disclosed here (SEQ ID NO: 1, 2, 3, and 34). SEQ ID NO: 3 includes the regions through the start of transcription and the start of translation, as evident from a sequence comparison to the figures. SEQ ID NO: 34 ends before the transcription start site, again as evident from a sequence comparison with the

figures. Six mutations are specifically disclosed here. TIGRmt1 is the result of a replacement of a cytosine with a guanine at position 4337 (SEQ ID NO: 1, SEQ ID NO: 2, and SEQ ID NO: 3). TIGRmt2 is the result of a replacement of a cytosine with a thymine at position 4950 (SEQ ID NO: 1, SEQ ID NO: 2, and SEQ ID NO: 3). TIGRmt3 is the result of an addition in the following order of a guanine, a thymine, a guanine, and a thymine (GTGT) at position 4998 (SEQ ID NO: 1, SEQ ID NO: 2, and SEQ ID NO: 3). TIGRmt4 is the result of a replacement of an adenine with a guanine at position 4256 (SEQ ID NO: 1, SEQ ID NO: 2, and SEQ ID NO: 3). TIGRmt5 is the result of a replacement of a guanine with an adenine at position 4262 (SEQ ID NO: 1, SEQ ID NO: 2 and SEQ ID NO: 3). TIGRmt11 (not pictured in Figure 2) is the result of a replacement of a thymine with a cytosine at position 5113 (SEQ ID NO: 1, 3, or 34) and at the equivalent position in SEQ ID NO: 2, at nucleotide 5117. One or more of TIGRmt1, TIGRmt2, TIGRmt3, TIGRmt4, TIGRmt5, and TIGTmt11 can be homozygous or heterozygous.

Sequence comparisons of the upstream region of a non-glaucoma individual and individuals with glaucoma identify at least one sequence variation in individuals with glaucoma. One such sequence variant is illustrated in Figure 2. *TIGRsv1* is the result of a replacement of an adenine with a guanine at position 4406 (SEQ ID NO: 1, SEQ ID NO: 2 and SEQ ID NO: 3). Also, the presence of *TIGRmt11* is associated with steroid sensitivity or an increased susceptibility to developing glaucoma or IOP-related disorders during steroid or corticosteroid treatment.

Molecules comprising sequences upstream of the TIGR coding region provide useful markers for polymorphic studies. Such molecules include primers suitable for single strand conformational polymorphic studies, examples of which are as follows: forward primer "Sk-1a": 5'-TGA GGC TTC CTC TGG AAA C-3' (SEQ ID NO: 6); reverse primer "ca2": 5'-TGA AAT CAG CAC ACC AGT AG-3' (SEQ ID NO: 7); forward primer "CA2": 5'-GCA CCC ATA CCC CAA TAA TAG-3' (SEQ ID NO: 8); reverse primer "Pr+1": 5'-AGA GTT CCC CAG ATT TCA CC-3' (SEQ ID NO: 9); forward primer "Pr-1": 5'-ATC TGG GGA ACT CTT CTC AG-3' (SEQ ID NO: 10); reverse primer "Pr+2(4A2)": 5'-TAC AGT TGT TGC AGA TAC G-3' (SEQ ID NO: 11); forward primer "Pr-2(4A)": 5'-ACA ACG TAT CTG CAA CAA CTG-3' (SEQ ID NO: 12); reverse primer "Pr+3(4A)": 5'-TCA GGC TTA ACT GCA GAA CC-3' (SEQ ID NO: 13); forward primer "Pr-3(4A)": 5'-TTG GTT CTG CAG TTA AGC C-3' (SEQ ID NO: 14); reverse primer "Pr+2(4A1)": 5'-AGC AGC ACA AGG GCA ATC C-3' (SEQ ID NO: 15); reverse primer "Pr+1(4A)": 5'-ACA GGG CTA TAT TGT GGG-3' (SEQ ID NO: 16).

In addition, molecules comprising sequences within TIGR exons provide useful markers for polymorphic studies. Such molecules include primers suitable for single strand conformational polymorphic studies, examples of which are as follows: forward primer "KS1X": 5'-CCT GAG ATG CCA GCT GTC C-3' (SEQ ID NO: 17); reverse primer "SK1XX": 5'-CTG

AAG CAT TAG AAG CCA AC-3' (SEQ ID NO: 18); forward primer "KS2a1": 5'-ACC TTG GAC CAG GCT GCC AG-3' (SEQ ID NO: 19); reverse primer "SK3" 5'-AGG TTT GTT CGA GTT CCA G-3' (SEQ ID NO: 20); forward primer "KS4": 5'-ACA ATT ACT GGC AAG TAT GG-3' (SEQ ID NO: 21); reverse primer "SK6A": 5'-CCT TCT CAG CCT TGC TAC C-3' (SEQ ID NO: 22); forward primer "KS5": 5'-ACA CCT CAG CAG ATG CTA CC-3' (SEQ ID NO: 23); reverse primer "SK8": 5'-ATG GAT GAC TGA CAT GGC C-3' (SEQ ID NO: 24); forward primer "KS6": 5'-AAG GAT GAA CAT GGT CAC C-3' (SEQ ID NO: 25).

The locations of primers: Sk-1a, ca2, CA2, Pr+1, Pr-1, Pr+2(4A2), Pr-2(4A), Pr+3(4A), Pr-3(4A), Pr-3(4A), Pr+2(4A1), and Pr+1(4A) are diagrammatically set forth in Figure 4. The location of primers: KS1X, SK1XX, Ks2a1, SK3, KS4, SK6A, KS5, SK8, and KS6 are diagrammatically set forth in Figure 5.

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The primary structure of the TIGR coding region initiates from an ATG initiation site (SEQ ID NO:3, residues 5337-5339) and includes a 20 amino acid consensus signal sequence a second ATG (SEQ ID NO: 3, residues 5379-5381), indicating that the protein is a secretory protein. The nucleotide sequence for the TIGR coding region is depicted in Figure 7 (SEQ ID NO: 26). The protein contains an N-linked glycosylation site located in the most hydrophilic region of the molecule. The amino terminal portion of the protein is highly polarized and adopts alpha helical structure as shown by its hydropathy profile and the Garnier-Robison structure analysis. In contrast, the protein contains a 25 amino acid hydrophobic region near its carboxy terminus. This region may comprise a glucocorticoid-induced protein (GIP) anchoring sequence. The amino acid sequence of TIGR is depicted in Figure 8 (SEQ ID NO: 32).

Study of cyclohexamide treatment in the absence and presence of GC suggest that the induction of TIGR may involve factors in addition to the GC receptor. The TIGR gene may be involved in the cellular stress response since it is also induced by stimulants such as H_2O_2 , 12-O-tetradecanolyphorbol-13-acetate (TPA), and high glucose; this fact may relate to glaucoma pathogenesis and treatment.

A preferred class of agents comprises TIGR nucleic acid molecules ("TIGR molecules") or fragments thereof. Such molecules may be either DNA or RNA. A second preferred class of agents ("TIGR molecules") comprises the TIGR protein, its peptide fragments, fusion proteins, and analogs.

TIGR nucleic acid molecules or fragments thereof are capable of specifically hybridizing to other nucleic acid molecules under certain circumstances. As used herein, two nucleic acid molecules are said to be capable of specifically hybridizing to one another if the two molecules are capable of forming an anti-parallel, double-stranded nucleic acid structure. A nucleic acid molecule is said to be the "complement" of another nucleic acid molecule if the molecules exhibit complete complementarity. As used herein, molecules are said to exhibit "complete

complementarity" when every nucleotide of one of the molecules is complementary to a nucleotide of the other. Two molecules are said to be "minimally complementary" if they can hybridize to one another with sufficient stability to permit them to remain annealed to one another under at least conventional "low-stringency" conditions. Similarly, the molecules are said to be "complementary" if they can hybridize to one another with sufficient stability to permit them to remain annealed to one another under conventional "high-stringency" conditions. Conventional stringency conditions are described by Sambrook et al., In: Molecular Cloning, A Laboratory Manual, 2nd Edition, Cold Spring Harbor Press, Cold Spring Harbor, New York (1989)), and by Haymes et al., In: Nucleic Acid Hybridization, A Practical Approach, IRL Press, Washington, DC (1985), the entirety of which is herein incorporated by reference. Departures from complete complementarity are therefore permissible, as long as such departures do not completely preclude the capacity of the molecules to form a double-stranded structure. In order for an nucleic acid molecule to serve as a primer or probe it need only be sufficiently complementary in sequence to be able to form a stable double-stranded structure under the particular solvent and salt concentrations employed.

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Appropriate stringency conditions which promote DNA hybridization, for example, 6.0 x sodium chloride/sodium citrate (SSC) at about 45°C, followed by a wash of 2.0 x SSC at 50°C, are known to those skilled in the art or can be found in Current Protocols in Molecular Biology, John Wiley & Sons, N.Y. (1989), 6.3.1-6.3.6. For example, the salt concentration in the wash step can be selected from a low stringency of about 2.0 x SSC at 50°C to a high stringency of about 0.2 x SSC at 50°C. In addition, the temperature in the wash step can be increased from low stringency conditions at room temperature, about 22°C, to high stringency conditions at about 65°C. Both temperature and salt may be varied, or either the temperature or the salt concentration may be held constant while the other variable is changed.

In a preferred embodiment, a nucleic acid of the present invention will specifically hybridize to one or more of the nucleic acid molecules set forth in SEQ ID NO: 1-5 or 34, or complements thereof, or fragments of about 20 to about 200 bases of either, under moderately stringent conditions, for example at about 2.0 x SSC and about 65°C. In a particularly preferred embodiment, a nucleic acid of the present invention will specifically hybridize to one or more of the nucleic acid molecules set forth in SEQ ID NO: 1-5 or 34, or complements or fragments of either under high stringency conditions.

In one aspect of the present invention, a preferred marker nucleic acid molecule of the present invention has the nucleic acid sequence set forth in SEQ ID NO: 6-25 or 33, or complements thereof or fragments of either. In another aspect of the present invention, a preferred marker nucleic acid molecule of the present invention shares between about 80% to about 100% or about 90% to about 100% sequence identity with the nucleic acid sequence set

forth in SEQ ID NO: 6-25 or 33, or complement thereof or fragments of either. In a further aspect of the present invention, a preferred marker nucleic acid molecule of the present invention shares between about 95% to about 100% sequence identity with the sequence set forth in SEQ ID NO: 6-25 or 33, or complement thereof or fragments of either. In a more preferred aspect of the present invention, a preferred marker nucleic acid molecule of the present invention shares between 98% and about 100% sequence identity with the nucleic acid sequence set forth in SEQ ID NO:6-25 or 33, or complement thereof or fragments of either.

Regulatory Regions and Agents that Bind to the Regions or Agents that Alter the Binding of a Molecule that Binds to the Regions

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Sequence comparisons of the upstream region identify a number of DNA motifs (cis elements) or regulatory regions. These DNA motifs or cis elements are shown in Figure 1. These motifs include, without limitation, glucocorticoid response motif(s), shear stress response motif(s), NFkB recognition motif(s), and AP1 motif(s). The locations of these and other motifs, discussed below, are diagramatically set forth in Figure 1.

As used herein, the term "cis elements capable of binding" refers to the ability of one or more of the described cis elements to specifically bind an agent. Such binding may be by any chemical, physical or biological interaction between the cis element and the agent, including, but not limited, to any covalent, steric, agostic, electronic and ionic interaction between the cis element and the agent. As used herein, the term "specifically binds" refers to the ability of the agent to bind to a specified cis element but not to wholly unrelated nucleic acid sequences.

Regulatory region refers to the ability of a nucleic acid fragment, region or length to functionally perform a biological activity. The biological activity may be binding to the nucleic or specific DNA sequence. The biological activity may also modulate, enhance, inhibit or alter the transcription of a nearby coding region. The biological activity may be identified by a gel shift assay, in which binding to a nucleic acid fragment can be detected. Other methods of detecting the biological activity in a nucleic acid regulatory region are known in the art (see Current Protocols in Molecular Biology, for example).

In an embodiment to identify or detect regulatory proteins or proteins or compounds that specifically bind to promoter or regulatory regions of the TIGR gene, a number of vector systems employing reporter genes can be useful. The promoter or 5' regulatory regions can be linked to control expression of the reporter gene so that the presence of the reporter gene indicates the transcriptional activity from the promoter or 5' regulatory region. The DNA constructs, vectors, and cells made for this aspect of the invention, and the methods employing them, can be useful in a variety of ways. For example, they can detect the presence or absence of tissue-specific factors

that modulate TIGR gene expression. Modulating expression can mean increasing or decreasing transcription, increasing or decreasing translation, or otherwise effecting the amount of TIGR mRNA or protein present in a cell. They can also be used to identify compounds that affect TIGR gene expression, IOP-related disorders, or steroid sensitivity. These identified compounds can then be used to development therapeutic or diagnostic treatments, especially for glaucoma and IOP-related disorders.

The types of expression systems and reporter genes that can be used or adapted for use are well known in the art. For example, vectors containing genes for a luciferase activity, an alkaline phosphatase activity, or a green fluorescent protein activity are commonly used. *See* Ausubel *et al.*, Current Protocols in Molecular Biology, John Wiley & Sons, N.Y. (1989), and supplements through May 1999. Inserting a 5' regulatory region of the TIGR gene into a vector of this type puts the expression of the reporter gene under the control of the TIGR sequence. The vector or vectors can then be inserted into various cell types to examine differences in tissue specific expression, which can then lead to identification of tissue specific factors that modulate TIGR expression as well as compounds that affect the activity of the tissue specific factors. The vectors can also be used to detect compounds that affect the transcription modulating activity of a TIGR 5' regulatory region. Alternatively, the 5' regulatory region and reporter gene constructs can be used in homologous recombination techniques to produce cells. When properly inserted into the genome of a cell, the reporter gene will be under the control of the TIGR regulatory region. Homologously recombinant cells can than be used, as discussed above, to detect changes in and modulation of expression due to various treatments or in separate cells.

For example, human trabecular meshwork cells (HTM) can be transiently or permanently transfected with a vector containing various 5' regulatory sequences from the TIGR gene that control expression of an alkaline phosphatase activity (AP). The expression of AP, or change in expression versus a control, indicates the presence of transcription-modulating sequences. By comparing the expression in HTM cells with the expression in other cells, such as Cos or HeLa cells, one can detect the presence of cell-specific compounds that influence transcriptional activity from the TIGR gene. Also, by treating the cells with test compounds, one can detect a compound that affects transcription.

Additionally, once cell types have been characterized for the presence of expression modulating activity, further uses of the 5' regulatory regions can identify the presence of specific DNA binding by cellular components or proteins within particular cells. Assays and methods for detection DNA binding include, but are not limited to, gel shift assays and equivalent techniques known in the art. In gel shift assays, the presence of a compound that binds to a DNA is detected by observing a shift in the mobility of the DNA through an agarose, polyacrylamide, or other gel matrix. Thus, by running the DNA through the gel in the presence of one or more cellular

components or cell extracts and comparing to controls, specific binding to the DNA can be detected. In a similar way, various compounds can be tested for their ability to affect binding of a cellular component to the DNA by treating the sample with one or more test compounds before running the DNA through the gel. Of course, the cells used in these assays need not be primary cultured cells or established cell lines. Any type of manipulated cell can be used, even those where deliberate changes to DNA binding proteins or transcription factors contained in the cell have been made.

The TIGR gene sequences that may be used in the DNA constructs, cells, vectors, or in methods for detecting useful promoter regions, tissue or cell-specific components that bind TIGR gene sequences, or in the related methods to detect compounds affecting DNA binding or promoter activity, can be large, for example at least as big as 1.6 kb, or much smaller, on the order of about 10 to 300 bp. A preferred region is the 283 bp region from 5340 to 5044 of SEQ ID NO: 3, designated SEQ ID NO: 37, or a 227 bp region from 5044 to 5271 (end) of SED ID NO: 34, which is identified as SEQ ID NO: 38.

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Other preferred regions are identified in the Figures and SEQ IDs listed or are inherently disclosed by comparing the Figures and SEQ IDs. To illustrate, the TATA boxes in SEQ ID NO: 3 and 34 both start at position 5232. Particular regions from SEQ ID NO: 3 and 34 can be correlated with the identified regulatory regions in Figure 1, where the TATA box begins at 5230. Thus, additional preferred 5' regions can incorporate one or more regulatory regions identified and discussed herein as well as the many sequence variations given in the consensus regulatory sequences listed in Figure 1. For example, SEQ ID NO: 37 can be changed at various places in the nGRE-Prl regulatory sequence, beginning at base 39 of SEQ ID NO: 37, to coincide with the consensus sequence given in Figure 1. Base 67 of SEQ ID NO: 37 can be changed from T to G, and bases 57 and 65 from C to A. These changes would not effect the basic regulatory activity of these sequences, but may affect the degree of regulatory modulation possible. Similar substitutions using any one or more of the listed regulatory regions can be made to a region from SEQ ID NO: 1-3, or 34. Also, substitutions that incorporate one or more of the mutant sequences identified herein may also be made. For example, base 69 of SEQ ID NO: 37 can be changed from a T to a C to correlate with the TIGR.mt11 mutant herein. Numerous variants in the 5' regulatory regions incorporating one or both of the consensus sequences changes and the changes from the identified TIGR mutants can be made. Bases may also be added or deleted without effecting the region's basic ability to modulate expression.

The following discussion identifies some of the regulatory regions identified in the TIGR 5'region or other well known regulatory acitivities. As noted, these regions or sequences, or variants of them, can be incorporated into the DNA constructs, vectors, cells, and methods of the invention. Any combination of them, with or without the above identified regions in SEQ ID

NO: 37 and 18, or variants thereof, can be tested for the ability to confer tissue specific, or cell-type specific expression of an attached gene. As shown in Example 5, below, SEQ ID NO: 37 confers tissue specific expression of an attached reporter gene. The expression of AP activity is much higher in the HTM cells than in HeLa or Cos cells. Thus, tissue specific promoter or regulatory activity can be identified by detecting a change of expression levels from one cell type compared to another.

Expression of the rat PRL gene is highly restricted to pituitary lactotroph cells and is induced by the cAMP-dependent protein kinase A pathway. At least one of the redundant pituitary specific elements (PRL-FP111) of the proximal rat PRL promotor is required for this protein kinase A effect (Rajnarayan et al., Molecular Endocrinology 4: 502-512 (1995), herein incorporated by reference). A sequence corresponding to an upstream motif or cis element characteristic of PRL-FP111 is set forth in Figure 1 at residues 370-388 and 4491-4502, respectively. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of molecules that bind the PRL-FP111 upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A consensus sequence (GR/PR), recognized by both the glucocorticoid receptor of rat liver and the progesterone receptor from rabbit uterus, has been reported to be involved in glucocorticoid and progesterone-dependent gene expression (Von der Ahe et al., Nature 313: 706-709 (1985), herein incorporated by reference). A sequence corresponding to a GC/PR upstream motif or cis element is set forth in Figure 1 at residues 433-445. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of glucocorticoid or progesterone or their homologues, including, but not limited to, the concentration of glucocorticoid or progesterone or their homologues bound to an GC/PR upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

Shear stress motif (SSRE) or cis element has been identified in a number of genes including platelet-derived growth factor B chain, tissue plasminogen activator (tPA), ICAM-1 and TGF-\(\beta\)1 (Resnick et al., Proc. Natl. Acad. Sci. (USA) 80: 4591-4595 (1993), herein incorporated by reference). Transcription of these genes has been associated with humoral stimuli such as cytokines and bacterial products as well as hemodynamic stress forces. Sequences corresponding to a upstream shear stress motif or cis element are set forth in Figure 1

at residues 446-451, 1288-1293, 3597-3602, 4771-4776, and 5240-5245, respectively. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of molecules capable of binding the shear stress motif. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A consensus sequence for a glucocorticoid response upstream motif (GRE) or cis element has been characterized (Beato, Cell 56: 335-344 (1989); Becker et al., Nature 324: 686-688 (1986), herein incorporated by reference; Sakai et al., Genes and Development 2: 1144-1154 (1988), herein incorporated by reference). Genes containing this upstream motif or cis element are regulated by glucocorticoids, progesterone, androgens and mineral corticoids (Beato, Cell 56: 335-344 (1989)). Sequences corresponding to glucocorticoid response upstream motif or cis element are set forth in Figure 1 at residues 574-600, 1042-1056, 2444-2468, 2442-2269, 3536-3563, 4574-4593, 4595-4614, 4851-4865, 4844-4864, 5079-5084, and 5083-5111, respectively. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of molecules capable of binding a glucocorticoid response upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

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A sequence specific binding site (CBE) for the wild type nuclear phosphoprotein, p53, has been identified and appears to be associated with replication origins (Kern et al. Science 252: 1708-1711 (1991), herein incorporated by reference). A sequence corresponding to an CBE upstream motif or cis element is set forth in Figure 1 at residues 735-746. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of p53 or its homologues, including, but not limited to, the concentration of p53 or its homologues bound to an CBE upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

Nuclear factor ets-like (NFE), a transcriptional activator that facilitates p50 and c-Rel-dependent IgH 3' enhancer activity has been shown to bind to an NFE site in the Rel-dependent IgH 3' enhancer (Linderson et al., European J. Immunology 27: 468-475 (1997), herein incorporated by reference). A sequence corresponding to an NFE upstream motif or cis element is set forth in Figure 1 at residues 774-795. In accordance with the embodiments of the present

invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of nuclear factors or their homologues, including, but not limited to, the concentration of nuclear factors or their homologues bound to an NFE upstream motif or *cis* element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

An upstream motif or cis element (KTF.1-CS) for a control element 3' to the human keratin 1 gene that regulates cell type and differentiation-specific expression has been identified (Huff et al., J. Biological Chemistry 268: 377-384 (1993), herein incorporated by reference). A sequence corresponding to an upstream motif or cis element characteristic of KTF.1-CS is set forth in Figure 1 at residues 843-854. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of KTF.1-CS or its homologues, including, but not limited to, the concentration of KTF.1-CS or its homologues bound to a KTF.1-CS upstream motif or cis element Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

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A progesterone responsive element (PRE) that maps to the far upstream steroid dependent DNase hypersensitive site of chicken lysozyme chromatin has been characterized (Hecht et al., EMBO J. 7: 2063-2073 (1988), herein incorporated by reference). The element confers hormonal regulation to a heterologous promoter and is composed of a cluster of progesterone receptor binding sites. A sequence corresponding to an upstream motif or cis element characteristic of PRE is set forth in Figure 1 at residues 987-1026. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of molecules capable of binding a progesterone responsive PRE upstream motif or cis element. Such agents may be useful in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A sequence (ETF-EGFR) has been characterized which serves as a motif for a transactive transcription factor that regulates expression of the epidermal growth factor receptor (Regec et al., Blood 85:2711-2719 (1995), herein incorporated by reference). A sequence corresponding to an ETF-EGFR upstream motif or cis element is set forth in Figure 1 at residues 1373-1388. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of nuclear factors or their homologues, including, but not limited to, the

concentration of nuclear factors or their homologues bound to an ETF-EGFR upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A common trans-acting factor (SRE-cFos) has been shown to regulate skeletal and cardiac alpha-Actin gene transcription in muscle (Muscat et al., Molecular and Cellular Biology 10: 4120-4133 (1988), herein incorporated by reference). A sequence corresponding to an SRE-cFos upstream motif or cis element is set forth in Figure 1 at residues 1447-1456. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of nuclear factors or their homologues, including, but not limited to, the concentration of nuclear factors or their homologues bound to an SRE-cFos upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

Alu repetitive elements are unique to primates and are interspersed within the human genome with an average spacing of 4Kb. While some Alu sequences are actively transcribed by polymerase III, normal transcripts may also contain Alu-derived sequences in 5' or 3' untranslated regions (Jurka and Mikahanljaia, J. Mol. Evolution 32: 105-121 (1991), herein incorporated by reference, Claveria and Makalowski, Nature 371: 751-752 (1994), herein incorporated by reference). A sequence corresponding to an Alu upstream motif or cis element is set forth in Figure 1 at residues 1331-1550. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of nuclear factors or their homologues, including, but not limited to, the concentration of nuclear factors or their homologues bound to an Alu upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A consensus sequence for a vitellogenin gene-binding protein (VBP) upstream motif or cis element has been characterized (Iyer et al., Molecular and Cellular Biology 11: 4863-4875 (1991), herein incorporated by reference). Expression of the VBP gene commences early in liver ontogeny and is not subject to circadian control. A sequence corresponding to an upstream motif or cis element capable of binding VBP is set forth in Figure 1 at residues 1786-1797. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of VBP or its homologues, including, but not limited to, the concentration of VBP or its homologues bound

to an VBP upstream motif or cis element Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A structural motif (Malt-CS) or cis element involved in the activation of all promoters of the maltose operons in Escherichia coli and Klebsiella pneumoniae has been characterized (Vidal-Ingigliardi et al., J. Mol. Biol. 218: 323-334 (1991), herein incorporated by reference). A sequence corresponding to a upstream Malt-CS motif or cis element is set forth in Figure 1 at residues 1832-1841. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of molecules capable of binding the upstream Malt-CS motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A consensus sequence for an estrogen receptor upstream motif or cis element has been characterized (ERE) (Forman et al., Mol. Endocrinology 4: 1293-1301 (1990), herein incorporated by reference; de Verneuil et al., Nucleic Acid Res. 18: 4489-4497 (1990), herein incorporated by reference; Gaub et al., Cell 63: 1267-1276 (1990), herein incorporated by reference. A sequence corresponding to half an upstream motif or cis element capable of binding estrogen receptor is set forth in Figure 1 at residues 2166-2195, 3413-3429, and 3892-3896, respectively. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration, of the estrogen receptor or its homologues bound to an upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

Certain protein-binding sites (NF-mutagen) in Ig gene enhancers which determine transcriptional activity and inducibility have been shown to interact with nuclear factors (Lenardo et al., Science 236: 1573-1577 (1987), herein incorporated by reference). A sequence corresponding to an NF-mutagen upstream motif or cis element is set forth in Figure 1 at residues 2329-2338. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of nuclear factors or their homologues, including, but not limited to, the concentration of nuclear factors or their homologues bound to an NF-mutagen upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A consensus sequence for a transcriptional repressor of c-myc (myc-PRF) upstream motif or cis element has been identified (Kakkis et al., Nature 339: 718-719 (1989), herein incorporated by reference). Myc-PRF interacts with another widely distributed protein, myc-CF1 (common factor 1), which binds nearby and this association may be important in myc-PRF repression. A sequence corresponding to an upstream motif or cis element capable of binding myc-PRF is set forth in Figure 1 at residues 2403-2416. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of myc-PRF or its homologues, including, but not limited to, the concentration of myc-PRF or its homologues bound to an myc-PRF upstream motif or cis element Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

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Human transcription factor activator protein 2 (AP2) is a transcription factor that has been shown to bind to Sp1, nuclear factor 1 (NF1) and simian virus 40 transplantation (SV40 T) antigen binding sites. It is developmentally regulated (Williams and Tijan, Gene Dev. 5: 670-682 (1991), herein incorporated by reference; Mitchell et al., Genes Dev. 5: 105-119 (1991), herein incorporated by reference; Coutois et al., Nucleic Acid Research 18: 57-64 (1990), herein incorporated by reference; Comb et al., Nucleic Acid Research 18: 3975-3982 (1990), herein incorporated by reference; Winings et al., Nucleic Acid Research 19: 3709-3714 (1991), herein Sequences corresponding to an upstream motif or cis element incorporated by reference). capable of binding AP2 are set forth in Figure 1 at residues 2520-2535, and 5170-5187, respectively. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of AP2 or its homologues, including, but not limited to, the concentration of AP2 or its homologues bound to an upstream motif or cis element. Such agents may be useful in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

Drosophila RNA polymerase II heat shock transcription factor (HSTF) is a transcription factor that has been shown to be required for active transcription of an hsp 70 gene (Parker and Topol, Cell 37: 273-283 (1984), herein incorporated by reference). Sequences corresponding to an upstream motif or cis element capable of binding HSTF are set forth in Figure 1 at residues 2622-2635, and 5105-5132. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of HSTF or its homologues, including, but not limited to, the concentration of HSTF or its homologues bound to an HSTF upstream motif or cis element.

Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A sequence corresponding to an upstream motif or cis element characteristic of SBF is set forth in Figure 1 at residues 2733-2743 (Shore et al., EMBO J. 6: 461-467 (1987), herein incorporated by reference). In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of molecules that bind the SBF upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

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An NF1 motif or cis element has been identified which recognizes a family of at least six proteins (Courtois, et al., Nucleic Acid Res. 18: 57-64 (1990), herein incorporated by reference; Mul et al., J. Virol. 64: 5510-5518 (1990), herein incorporated by reference; Rossi et al., Cell 52: 405-414 (1988), herein incorporated by reference; Gounari et al., EMBO J. 10: 559-566 (1990), herein incorporated by reference; Goyal et al., Mol. Cell Biol. 10: 1041-1048 (1990); herein incorporated by reference; Mermond et al., Nature 332: 557-561 (1988), herein incorporated by reference; Gronostajski et al., Molecular and Cellular Biology 5: 964-971 (1985), herein incorporated by reference; Hennighausen et al., EMBO J. 5: 1367-1371 (1986), herein incorporated by reference; Chodosh et al., Cell 53: 11-24 (1988), herein incorporated by reference). The NF1 protein will bind to an NF1 motif or cis element either as a dimer (if the motif is palindromic) or as an single molecule (if the motif is not palindromic). The NF1 protein is induced by TGFB (Faisst and Meyer, Nucleic Acid Research 20: 3-26 (1992), herein incorporated by reference). Sequences corresponding to an upstream motif or cis element capable of binding NF1 are set forth in Figure 1 at residues 2923-2938, 4143-4167, and 4886-4900, respectively. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of NF1 or its homologues, including, but not limited to, the concentration of NF1 or its homologues bound to an upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

Conserved regulatory sequences (NF-MHCIIA/B) of a rabbit major histocompatability complex (MHC) class II gene are responsible for binding two distinct nuclear factors NF-MHCIIA and NF-MHCIIB and are believed to be involved in the regulation of coordinate expression of the class II genes — eg. MHC class II gene in B lymphocytes (Sittisombut

Molecular and Cellular Biology 5: 2034-2041 (1988), herein incorporated by reference). A sequence corresponding to an NF-MHCIIA/B upstream motif or cis element is set forth in Figure 1 at residues 2936-2944. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of NF-MHCIIA or NF-MHCIIB or their homologues, including, but not limited to, the concentration of NF-MHCIIA or NF-MHCIIB or their homologues bound to an NF-MHCIIA/B upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

PEA 1 binding motifs or cis elements have been identified (Piette and Yaniv, EMBO J. 5: 1331-1337 (1987), herein incorporated by reference). The PEA1 protein is a transcription factor that is reported to bind to both the polyoma virus and c-fos enhancers. A sequence corresponding to an upstream motif or cis element capable of binding PEA1 is set forth in Figure 1 at residues 3285-3298. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of PEA1 or its homologues, including, but not limited to, the concentration of PEA1 or its homologues bound to an upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A conserved cis-acting regulatory element (ICS) has been shown to bind trans-acting constitutive nuclear factors present in lymphocytes and fibroblasts which are involved in the interferon (IFN)-mediated transcriptional enhancement of MHC class I and other genes (Shirayoshi et al., Proc. Natl. Acad. Sci. (USA) 85: 5884-5888 (1988), herein incorporated by reference). A sequence corresponding to an ICS upstream motif or cis element is set forth in Figure 1 at residues 3688-3699. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of nuclear factors or their homologues, including, but not limited to, the concentration of nuclear factors or their homologues bound to an ICS upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A consensus sequence for an ISGF2 upstream motif or cis element has been characterized (Iman et al., Nucleic Acids Res. 18: 6573-6580 (1990), herein incorporated by reference; Harada et al., Cell 63: 303-312 (1990), herein incorporated by reference; Yu-Lee et al., Mol. Cell Biol.

10: 3087-3094 (1990), herein incorporated by reference; Pine et al., Mol. Cell Biol. 10: 32448-2457 (1990), herein incorporated by reference). ISGF2 is induced by interferon α and γ , prolactin and virus infections. A sequence corresponding to an upstream motif or cis element capable of binding ISGF2 is set forth in Figure 1 at residues 4170-4179. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of ISGF2 or its homologues, including, but not limited to, the concentration of ISGF2 or its homologues bound to an upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A sequence corresponding to an upstream motif or *cis* element capable of binding zinc is set forth in Figure 1 at residues 4285-4292. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of zinc. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A sequence corresponding to an upstream motif or *cis* element characteristic of CAP/CRP-galO is set forth in Figure 1 at residues 4379-4404 (Taniguchi *et al.*, *Proc. Natl. Acad. Sci (USA) 76:* 5090-5094 (1979), herein incorporated by reference). In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of molecules that bind the CAP/CRP-galO upstream motif or *cis* element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

Human transcription factor activator protein 1 (AP1) is a transcription factor that has been shown to regulate genes which are highly expressed in transformed cells such as stromelysin, c-fos, α₁-anti-trypsin and collagenase (Gutman and Wasylyk, EMBO J. 9.7: 2241-2246 (1990), herein incorporated by reference; Martin et al., Proc. Natl. Acad. Sci. USA 85: 5839-5843 (1988), herein incorporated by reference; Jones et al., Genes and Development 2: 267-281 (1988), herein incorporated by reference; Faisst and Meyer, Nucleic Acid Research 20: 3-26 (1992), herein incorporated by reference; Kim et al., Molecular and Cellular Biology 10: 1492-1497 (1990), herein incorporated by reference: Baumhueter et al., EMBO J. 7: 2485-2493 (1988), herein incorporated by reference). The AP1 transcription factor has been associated with genes that are activated by 12-O-tetradecanolyphorbol-13-acetate (TPA) (Gutman and Wasylyk,

EMBO J.7: 2241-2246 (1990)). Sequences corresponding to an upstream motif or cis element capable of binding AP1 are set forth in Figure 1 at residues 4428-4434 and 4627-4639, respectively. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of AP1 or its homologues, including, but not limited to, the concentration of AP1 or its homologues bound to an upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

The sex-determining region of the Y chromosome gene, sry, is expressed in the fetal mouse for a brief period, just prior to testis differentiation. SRY is a DNA binding protein known to bind to a CACA-rich region in the sry gene (Vriz et al., Biochemistry and Molecular Biology International 37: 1137-1146 (1995), herein incorporated by reference). A sequence corresponding to an upstream motif or cis element capable of binding SRY is set forth in Figure 1 at residues 4625-4634. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of SRY or its homologues, including, but not limited to, the concentration of SRY or its homologues bound to an upstream motif or cis element. Such agents may be useful in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A sequence corresponding to an upstream motif or cis element characteristic of GC2-GH is set forth in Figure 1 at residues 4689-4711 (West et al., Molecular and Cellular Biology 7: 1193-1197 (1987), herein incorporated by reference). In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of GC2-GH or its homologues, including, but not limited to, the concentration of GC2-GH or its homologues bound to an upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

PEA 3 binding motifs or cis elements have been identified (Martin et al., Proc. Natl. Acad. Sci. (USA) 85: 5839-5843 (1988), herein incorporated by reference; Gutman and Wasylyk, EMBO J. 7: 2241-2246 (1990), herein incorporated by reference). The PEA3 protein is a transcription factor that is reported to interact with AP1 like proteins (Martin et al., Proc. Natl. Acad. Sci. (USA) 85: 5839-5843 (1988), herein incorporated by reference). Sequences corresponding to an upstream motif or cis element capable of binding PEA3 is set forth in Figure

1 at residues 4765-4769. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of PEA3 or its homologues, including, but not limited to, the concentration of PEA3 or its homologues bound to an upstream motif or *cis* element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

Mammalian interspersed repetitive (MIR) is an element involved in the coding and processing sequences of mammalian genes. The MIR element is at least 260 bp in length and numbers about 10⁵ copies within the mammalian genome (Murnane et al., Nucleic Acids Research 15: 2837-2839 (1995), herein incorporated by reference). A sequence corresponding to an MIR upstream motif or cis element is set forth in Figure 1 at residues 4759-4954. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of nuclear factors or their homologues, including, but not limited to, the concentration of nuclear factors or their homologues bound to an MIR upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

Normal liver and differentiated hepatoma cell lines contain a hepatocyte-specific nuclear factor (HNF-1) which binds cis-acting element sequences within the promoters of the alpha and beta chains of fibrinogen and alpha 1-antitrypsin (Baumhueter et al., EMBO J. 8: 2485-2493, herein incorporated by reference). A sequence corresponding to an HNF-1 upstream motif or cis element is set forth in Figure 1 at residues 4923-4941. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of HNF-1 or its homologues, including, but not limited to, the concentration of HNF-1 or its homologues bound to an HNF-1 upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

A number of cis elements or upstream motifs have been associated with gene regulation by steroid and thyroid hormones (e.g. glucocorticoid and estrogen)(Beato, Cell 56: 335-344 (1989), herein incorporated by reference; Brent et al., Molecular Endocrinology 89:1996-2000 (1989), herein incorporated by reference; Glass et al., Cell 54: 313-323 (1988), herein incorporated by reference; Evans, Science 240: 889-895 (1988), herein incorporated by reference).

A consensus sequence for a thyroid receptor upstream motif or *cis* element (TRE) has been characterized (Beato, *Cell 56*: 335-344 (1989), herein incorporated by reference). A sequence corresponding to a thyroid receptor upstream motif or *cis* element is set forth in Figure 1 at residues 5151-5156. Thyroid hormones are capable of regulating genes containing a thyroid receptor upstream motif or *cis* element (Glass *et al.*, *Cell 54*: 313-323 (1988), herein incorporated by reference). Thyroid hormones can negatively regulate TIGR. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of molecules capable of binding a thyroid receptor upstream motif or *cis* element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

NFkB is a transcription factor that is reportedly associated with a number of biological processes including T-cell activation and cytokine regulation (Lenardo et al., Cell 58: 227-229 (1989), herein incorporated by reference). A consensus upstream motif or cis element capable of binding NFkB has been reported (Lenardo et al., Cell 58: 227-229 (1989)). Sequences corresponding to an upstream motif or cis element capable of binding NFkB are set forth in Figure 1 at residues 5166-5175. In accordance with the embodiments of the present invention, transcription of TIGR molecules can be effected by agents capable of altering the biochemical properties or concentration of NFkB or its homologues, including, but not limited to, the concentration of NFkB or its homologues bound to an upstream motif or cis element. Such agents can be used in the study of glaucoma pathogenesis. In another embodiment, such agents can also be used in the study of glaucoma prognosis. In another embodiment such agents can be used in the treatment of glaucoma.

25 Illustrative Uses of the Nucleic Acids of the Invention

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Where one or more of the agents is a nucleic acid molecule, such nucleic acid molecule may be sense, antisense or triplex oligonucleotides corresponding to any part of the TIGR promoter, TIGR cDNA, TIGR intron, TIGR exon or TIGR gene. In some embodiments these nucleic acids may be about 20 bases in length, as for example, SEQ. ID NO: 6-25 or 33. In some circmstances, the nucleic acids may be only about 8 bases in length. Short nucleic acids may be particularly useful in hybridization to immobilized nucleic acids in order to determine the presence of specific sequences, such as by the known methods of sequencing by hybridization.

The TIGR promoter, or fragment thereof, of the present invention may be cloned into a suitable vector and utilized to promote the expression of a marker gene (e.g. firefly luciferase (de

Wet, Mol. Cell Biol. 7: 725-737 (1987), herein incorporated by reference) or GUS (Jefferson et al., EMBO J. 6: 3901-3907 (1987), herein incorporated by reference)).

In another embodiment of the present invention, a TIGR promoter may be cloned into a suitable vector and utilized to promote the expression of a TIGR gene in a suitable eukaryotic or prokaryotic host cell (e.g. human trabecular cell, chinese hamster cell, *E. coli*). In another embodiment of the present invention, a TIGR promoter may be cloned into a suitable vector and utilized to promote the expression of a homologous or heterologous gene in a suitable eukaryotic or prokaryotic host cells (e.g. human trabecular cell lines, chinese hamster cells, *E. coli*).

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Similarly, the TIGR promoter, regions of the TIGR 5' regulatory region from about 10 bp to about 1.6 kb, or regions of the TIGR 5' regulatory region that confer tissue specific expression can be used in DNA constructs, vectors, and cells to express an operably linked gene. For example, these TIGR sequences can control expression of a linked gene in a vector used for producing transgenic animals. In this way, cells or animals can be produced that express a gene in a tissue specific manner. One embodiment involves a gene therapy approach to treating glaucoma or IOP-disorder, where a transgene having the TIGR 5'regulatory sequence is used to direct expression of a therapeutic gene into the trabecular meshwork or the cells of the trabecular meshwork. Experimental animal models can also be produced using the transgenic technology. These animal models, or cells from them or comprising these TIGR sequences, can be used in methods to screen for compounds that modulate expression of TIGR, and especially compounds that affect the steroid regulation of TIGR expression. These methods will produce therapeutic or diagnostic agents, which agents are specifically included in the invention.

Thus, transgenic animals having an introduced DNA that comprises at least a portion of the TIGR 5' regulatory region are specifically included in this invention. Preferred embodiments have 5' regulatory sequences of SEQ ID NO: 37 or 38, or variant thereof, or a region of SEQ ID NO: 3 or 34, especially those that confer tissue specific expression. One skilled in the art is familiar with the production of transgenic construct, vectors, and animals (See, Ausubel et al., Current Protocols in Molecular Biology).

In addition, as noted above, the 5' regulatory regions can be used to screen and identify DNA binding proteins. The specific binding of proteins or cellular components to the 5' regulatory regions can be detected, see Example 5. Screening methods for compounds that modulate the binding can also be produced using the 5'regulatory regions of the invention, or vectors or cells comprising those regions. DNA binding proteins that increase or decrease expression can be identified. For example, cells that do not show native expression of TIGR, such as HeLa cells, may possess proteins or cellular components that inhibit expression of TIGR. While cells like TM cells may possess DNA binding proteins or cellular components that allow expression of TIGR or increase the level of TIGR expression in response to other agents, such as

steroids or dexamethasone. Thus, the invention encompasses a number of methods to detect and identify proteins or cellular components that modulate expression of TIGR, or the expression of a gene under the control of the TIGR 5' regulatory region. Also, methods to detect compounds that effect that modulation are similarly encompassed, as well as therapeutic and diagnostic agents that are identified from those methods.

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Practitioners are familiar with the standard resource materials which describe specific conditions and procedures for the construction, manipulation and isolation of macromolecules (e.g., DNA molecules, plasmids, etc.), generation of recombinant organisms and the screening and isolating of clones, (see for example, Sambrook et al., In Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Press (1989), herein incorporated by reference in its entirety; Old and Primrose, In Principles of Gene Manipulation: An Introduction To Genetic Engineering, Blackwell (1994), herein incorporated by reference).

The TIGR promoter, or any portion thereof, or an about 10 to about 500 bases fragment thereof, of the present invention may be used in a gel-retardation or band shift assay (Old and Primrose, In Principles of Gene Manipulation: An Introduction To Genetic Engineering, Blackwell (1994)). Nucleic acids or fragments comprising any of the cis elements identified in the present invention may be used in a gel-retardation or band shift assay to isolate proteins capable of binding the cis element. Suitable DNA fragments or molecules comprise or consist of one or more of the following: sequences corresponding to an upstream motif or cis element characteristic of PRL-FP111 as set forth in Figure 1 at residues 370-388, and 4491-4502, respectively, a sequence corresponding to an upstream motif or cis element capable of binding GR/PR as set forth in Figure 1 at residues 433-445, sequences corresponding to an upstream shear stress motif or cis element as set forth in Figure 1 at residues 446-451, 1288-1293, 3597-3602, 4771-4776, and 5240-5245, respectively, sequences corresponding to glucocorticoid response upstream motif or cis element as set forth in Figure 1 at residues 574-600, 1042-1056, 2444-2468, 2442-2269, 3536-3563, 4574-4593, 4595-4614, 4851-4865, 4844-4864, 5079-5084, 5083-5111, respectively, a sequence corresponding to an upstream motif or cis element capable of binding CBE as set forth in Figure 1 at residues 735-746, a sequence corresponding to an upstream motif or cis element capable of binding NFE as set forth in Figure 1 at residues 774-795, a sequence corresponding to an upstream motif or cis element capable of binding KTF.1-CS as set forth in Figure 1 at residues 843-854, a sequence corresponding to an upstream motif or cis element capable of binding PRE is set forth in Figure 1 at residues 987-1026, a sequence corresponding to an upstream motif or cis element capable of binding ETF-EGFR as set forth in Figure 1 at residues 1373-1388, a sequence corresponding to an upstream motif or cis element capable of binding SRE-cFos as set forth in Figure 1 at residues 1447-1456, a sequence corresponding to an upstream motif or cis element capable of binding Alu as set forth in Figure

1 at residues 1331-1550, a sequence corresponding to an upstream motif or cis element capable of binding VBP as set forth in Figure 1 at residues 1786-1797, a sequence corresponding to an upstream motif or cis element capable of binding Malt-CS as set forth in Figure 1 at residues 1832-1841, sequences corresponding to an upstream motif or cis element capable of binding ERE as set forth in Figure 1 at residues 2167-2195, 3413-3429, and 3892-3896, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-mutagen as set forth in Figure 1 at residues 2329-2338, a sequence corresponding to an upstream motif or cis element capable of binding myc-PRF as set forth in Figure 1 at residues 2403-2416, sequences corresponding to an upstream motif or cis element capable of binding AP2 as set forth in Figure 1 at residues 2520-2535 and 5170-5187, respectively, sequences corresponding to an upstream 10 motif or cis element capable of binding HSTF as set forth in Figure 1 at residues 2622-2635, and 5105-5132, respectively, a sequence corresponding to an upstream motif or cis element characteristic of SBF as set forth in Figure 1 at residues 2733-2743, sequences corresponding to an upstream motif or cis element capable of binding NF-1 as set forth in Figure 1 at residues 2923-2938, 4144-4157, and 4887-4900, respectively, a sequence corresponding to an upstream 15 motif or cis element capable of binding NF-MHCIIA/B as set forth in Figure 1 at residues 2936-2944, a sequence corresponding to an upstream motif or cis element capable of binding PEA1 as set forth in Figure 1 at residues 3285-3298, a sequence corresponding to an upstream motif or cis element capable of binding ICS as set forth in Figure 1 at residues 3688-3699, a sequence corresponding to an upstream motif or cis element capable of binding ISGF2 as set forth in 20 Figure 1 at residues 4170-4179, a sequence corresponding to an upstream motif or cis element capable of binding zinc as set forth in Figure 1 at residues 4285-4293, a sequence corresponding to an upstream motif or cis element characteristic of CAP/CRP-galO as set forth in Figure 1 at residues 4379-4404, sequences corresponding to an upstream motif or cis element capable of binding AP1 as set forth in Figure 1 at residues 4428-4434, and 4627-4639, respectively, a 25 sequence corresponding to an upstream motif or cis element capable of binding SRY as set forth in Figure 1 at residues 4625-4634, a sequence corresponding to an upstream motif or cis element characteristic of GC2 as set forth in Figure 1 at residues 4678-4711, a sequence corresponding to an upstream motif or cis element capable of binding PEA3 as set forth in Figure 1 at residues 4765-4769, a sequence corresponding to an upstream motif or cis element capable of MIR as set 30 forth in Figure 1 at residues 4759-4954, a sequence corresponding to an upstream motif or cis element capable of binding NF-HNF-1 as set forth in Figure 1 at residues 4923-4941, a sequence corresponding to a thyroid receptor upstream motif or cis element as set forth in Figure 1 at residues 5151-5156, and a sequence corresponding to an upstream motif or cis element capable of binding NFkB as set forth in Figure 1 at residues 5166-5175. 35

A preferred class of agents of the present invention comprises nucleic acid molecules encompassing all or a fragment of the "TIGR promoter" or 5' flanking gene sequences. As used herein, the terms "TIGR promoter" or "promoter" is used in an expansive sense to refer to the regulatory sequence(s) that control mRNA production. Thus, TIGR promoter sequences can be identified by those sequences that functionally effect the intiation, rate, or amount of transcription of the TIGR gene product mRNA. Such sequences include RNA polymerase binding sites, glucocorticoid response elements, enhancers, etc. These sequences may preferably be found within the specifically disclosed 5' upstream region sequences disclosed here, and most preferably within an about 500 base region 5' to the start of transcription or within an about 300 base region 5' of the transcription start site. However, other genomic sequences may be a TIGR promoter. Methods known in the art to identify distant promoter elements can be used with the disclosed sequences and nucleic acids to identify and define these distant TIGR promoter sequences. Such TIGR molecules may be used to diagnose the presence of glaucoma and the severity of or susceptibility to glaucoma. Such molecules may be either DNA or RNA.

A functional regulatory region of the TIGR gene may be a TIGR promoter sequence. It may also include transcription enhancer sites and transcription inhibitor sites or binding sites for a number of known proteins or molecules demonstrated as effecting transcription. A number of regulatory elements are discussed below, and the equivalent of those activities can represent the functional regulatory region of the TIGR gene. The methods for identifying and detecting the activity and function of these regulatory regions are known in the art.

Fragment nucleic acid molecules may encode significant portion(s) of, or indeed most of, SEQ ID NO: 1 or SEQ ID NO: 3 or SEQ ID NO: 4 or SEQ ID NO: 5. Alternatively, the fragments may comprise smaller oligonucleotides (having from about 15 to about 250 nucleotide residues, and more preferably, about 15 to about 30 nucleotide residues.). Such oligonucleotides include SEQ ID NO: 6, SEQ ID NO: 7, SEQ ID NO: 8, SEQ ID NO: 9, SEQ ID NO: 10, SEQ ID NO: 11, SEQ ID NO: 12, SEQ ID NO: 13, SEQ ID NO: 14, SEQ ID NO: 15, SEQ ID NO: 16, SEQ ID NO: 17, SEQ ID NO: 18, SEQ ID NO: 19, SEQ ID NO: 20, SEQ ID NO: 21, SEQ ID NO: 22, SEQ ID NO: 23, SEQ ID NO: 24, SEQ ID NO: 25.

Alternatively such oligonucleotides may derive from either the TIGR promoter, TIGR introns, TIGR exons, TIGR cDNA and TIGR downstream sequences comprise or consist of one or more of the following: sequences corresponding to an upstream motif or *cis* element characteristic of PRL-FP111 as set forth in Figure 1 at residues 370-388, and 4491-4502, respectively, a sequence corresponding to an upstream motif or *cis* element capable of binding GR/PR as set forth in Figure 1 at residues 433-445, sequences corresponding to an upstream shear stress motif or *cis* element as set forth in Figure 1 at residues 446-451, 1288-1293, 3597-3602, 4771-4776, and 5240-5245, respectively, sequences corresponding to glucocorticoid

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response upstream motif or cis element as set forth in Figure 1 at residues 574-600, 1042-1056, 2444-2468, 2442-2269, 3536-3563, 4574-4593, 4595-4614, 4851-4865, 4844-4864, 5079-5084, 5083-5111, respectively, a sequence corresponding to an upstream motif or cis element capable of binding CBE as set forth in Figure 1 at residues 735-746, a sequence corresponding to an upstream motif or cis element capable of binding NFE as set forth in Figure 1 at residues 774-795, a sequence corresponding to an upstream motif or cis element capable of binding KTF.1-CS as set forth in Figure 1 at residues 843-854, a sequence corresponding to an upstream motif or cis element capable of binding PRE is set forth in Figure 1 at residues 987-1026, a sequence corresponding to an upstream motif or cis element capable of binding ETF-EGFR as set forth in Figure 1 at residues 1373-1388, a sequence corresponding to an upstream motif or cis element capable of binding SRE-cFos as set forth in Figure 1 at residues 1447-1456, a sequence corresponding to an upstream motif or cis element capable of binding Alu as set forth in Figure 1 at residues 1331-1550, a sequence corresponding to an upstream motif or cis element capable of binding VBP as set forth in Figure 1 at residues 1786-1797, a sequence corresponding to an upstream motif or cis element capable of binding Malt-CS as set forth in Figure 1 at residues 1832-1841, sequences corresponding to an upstream motif or cis element capable of binding ERE as set forth in Figure 1 at residues 2167-2195, 3413-3429, and 3892-3896, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-mutagen as set forth in Figure 1 at residues 2329-2338, a sequence corresponding to an upstream motif or cis element capable of binding myc-PRF as set forth in Figure 1 at residues 2403-2416, sequences corresponding to an upstream motif or cis element capable of binding AP2 as set forth in Figure 1 at residues 2520-2535 and 5170-5187, respectively, sequences corresponding to an upstream motif or cis element capable of binding HSTF as set forth in Figure 1 at residues 2622-2635, and 5105-5132, respectively, a sequence corresponding to an upstream motif or cis element characteristic of SBF as set forth in Figure 1 at residues 2733-2743, sequences corresponding to an upstream motif or cis element capable of binding NF-1 as set forth in Figure 1 at residues 2923-2938, 4144-4157, and 4887-4900, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-MHCIIA/B as set forth in Figure 1 at residues 2936-2944, a sequence corresponding to an upstream motif or cis element capable of binding PEA1 as set forth in Figure 1 at residues 3285-3298, a sequence corresponding to an upstream motif or cis 30 element capable of binding ICS as set forth in Figure 1 at residues 3688-3699, a sequence corresponding to an upstream motif or cis element capable of binding ISGF2 as set forth in Figure 1 at residues 4170-4179, a sequence corresponding to an upstream motif or cis element capable of binding zinc as set forth in Figure 1 at residues 4285-4293, a sequence corresponding to an upstream motif or cis element characteristic of CAP/CRP-galO as set forth in Figure 1 at 35 residues 4379-4404, sequences corresponding to an upstream motif or cis element capable of

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binding AP1 as set forth in Figure 1 at residues 4428-4434, and 4627-4639, respectively, a sequence corresponding to an upstream motif or *cis* element capable of binding SRY as set forth in Figure 1 at residues 4625-4634, a sequence corresponding to an upstream motif or *cis* element characteristic of GC2 as set forth in Figure 1 at residues 4678-4711, a sequence corresponding to an upstream motif or *cis* element capable of binding PEA3 as set forth in Figure 1 at residues 4765-4769, a sequence corresponding to an upstream motif or *cis* element capable of MIR as set forth in Figure 1 at residues 4759-4954, a sequence corresponding to an upstream motif or *cis* element capable of binding NF-HNF-1 as set forth in Figure 1 at residues 4923-4941, a sequence corresponding to a thyroid receptor upstream motif or *cis* element as set forth in Figure 1 at residues 5151-5156, and a sequence corresponding to an upstream motif or *cis* element capable of binding NFkB as set forth in Figure 1 at residues 5166-5175. For such purpose, the oligonucleotides must be capable of specifically hybridizing to a nucleic acid molecule genetically or physically linked to the TIGR gene. As used herein, the term "linked" refers to genetically, physically or operably linked.

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As used herein, two nucleic acid molecules are said to be capable of specifically hybridizing to one another if the two molecules are capable of forming an anti-parallel, doublestranded nucleic acid structure, whereas they are unable to form a double-stranded structure when incubated with a non-TIGR nucleic acid molecule. A nucleic acid molecule is said to be the "complement" of another nucleic acid molecule if they exhibit complete complementarity. As used herein, molecules are said to exhibit "complete complementarity" when every nucleotide of one of the molecules is complementary to a nucleotide of the other. Two molecules are said to be "minimally complementary" if they can hybridize to one another with sufficient stability to permit them to remain annealed to one another under at least conventional "low-stringency" conditions. Similarly, the molecules are said to be "complementary" if they can hybridize to one another with sufficient stability to permit them to remain annealed to one another under conventional "high-stringency" conditions. Conventional stringency conditions are described by Sambrook, J., et al., (In: Molecular Cloning, a Laboratory Manual, 2nd Edition, Cold Spring Harbor Press, Cold Spring Harbor, New York (1989)), and by Haymes, B.D., et al. (In: Nucleic Acid Hybridization, A Practical Approach, IRL Press, Washington, DC (1985)), both herein incorporated by reference). Departures from complete complementarity are therefore permissible, as long as such departures do not completely preclude the capacity of the molecules to form a double-stranded structure. Thus, in order for an oligonucleotide to serve as a primer it need only be sufficiently complementary in sequence to be able to form a stable double-stranded structure under the particular solvent and salt concentrations employed.

Apart from their diagnostic or prognostic uses, such oligonucleotides may be employed to obtain other TIGR nucleic acid molecules. Such molecules include the TIGR-encoding nucleic

acid molecule of non-human animals (particularly, cats, monkeys, rodents and dogs), fragments thereof, as well as their promoters and flanking sequences. Such molecules can be readily obtained by using the above-described primers to screen cDNA or genomic libraries obtained from non-human species. Methods for forming such libraries are well known in the art. Such analogs may differ in their nucleotide sequences from that of SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7, SEQ ID NO: 8, SEQ ID NO: 9, SEQ ID NO: 10, SEQ ID NO: 11, SEQ ID NO: 12, SEQ ID NO: 13, SEQ ID NO: 14, SEQ ID NO: 15, SEQ ID NO: 16, SEQ ID NO: 17, SEQ ID NO: 18, SEQ ID NO: 19, SEQ ID NO: 20, SEQ ID NO: 21, SEQ ID NO: 22, SEQ ID NO: 23, SEQ ID NO: 24, SEQ ID NO: 25, or from molecules consisting of sequences corresponding to an upstream motif or cis element characteristic of PRL-FP111 as set forth in Figure 1 at residues 370-388, and 4491-4502, respectively, a sequence corresponding to an upstream motif or cis element capable of binding GR/PR as set forth in Figure 1 at residues 433-445, sequences corresponding to an upstream shear stress motif or cis element as set forth in Figure 1 at residues 446-451, 1288-1293, 3597-3602, 4771-4776, and 5240-5245, respectively, sequences corresponding to glucocorticoid response upstream motif or cis element as set forth in Figure 1 at residues 574-600, 1042-1056, 2444-2468, 2442-2269, 3536-3563, 4574-4593, 4595-4614, 4851-4865, 4844-4864, 5079-5084, 5083-5111, respectively, a sequence corresponding to an upstream motif or cis element capable of binding CBE as set forth in Figure 1 at residues 735-746, a sequence corresponding to an upstream motif or cis element capable of binding NFE as set forth in Figure 1 at residues 774-795, a sequence corresponding to an upstream motif or cis element capable of binding KTF.1-CS as set forth in Figure 1 at residues 843-854, a sequence corresponding to an upstream motif or cis element capable of binding PRE is set forth in Figure 1 at residues 987-1026, a sequence corresponding to an upstream motif or cis element capable of binding ETF-EGFR as set forth in Figure 1 at residues 1373-1388, a sequence corresponding to an upstream motif or cis element capable of binding SRE-cFos as set forth in Figure 1 at residues 1447-1456, a sequence corresponding to an upstream motif or cis element capable of binding Alu as set forth in Figure 1 at residues 1331-1550, a sequence corresponding to an upstream motif or cis element capable of binding VBP as set forth in Figure 1 at residues 1786-1797, a sequence corresponding to an upstream motif or cis element capable of binding Malt-CS as set forth in Figure 1 at residues 1832-1841, sequences corresponding to an upstream motif or cis element capable of binding ERE as set forth in Figure 1 at residues 2167-2195, 3413-3429, and 3892-3896, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-mutagen as set forth in Figure 1 at residues 2329-2338, a sequence corresponding to an upstream motif or cis element capable of binding myc-PRF as set forth in Figure 1 at residues 2403-2416, sequences corresponding to an upstream motif or cis element capable of binding AP2 as set forth in Figure

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1 at residues 2520-2535 and 5170-5187, respectively, sequences corresponding to an upstream motif or cis element capable of binding HSTF as set forth in Figure 1 at residues 2622-2635, and 5105-5132, respectively, a sequence corresponding to an upstream motif or cis element characteristic of SBF as set forth in Figure 1 at residues 2733-2743, sequences corresponding to an upstream motif or cis element capable of binding NF-1 as set forth in Figure 1 at residues 2923-2938, 4144-4157, and 4887-4900, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-MHCIIA/B as set forth in Figure 1 at residues 2936-2944, a sequence corresponding to an upstream motif or cis element capable of binding PEA1 as set forth in Figure 1 at residues 3285-3298, a sequence corresponding to an upstream motif or cis element capable of binding ICS as set forth in Figure 1 at residues 3688-3699, a sequence corresponding to an upstream motif or cis element capable of binding ISGF2 as set forth in Figure 1 at residues 4170-4179, a sequence corresponding to an upstream motif or cis element capable of binding zinc as set forth in Figure 1 at residues 4285-4293, a sequence corresponding to an upstream motif or cis element characteristic of CAP/CRP-galO as set forth in Figure 1 at residues 4379-4404, sequences corresponding to an upstream motif or cis element capable of binding AP1 as set forth in Figure 1 at residues 4428-4434, and 4627-4639, respectively, a sequence corresponding to an upstream motif or cis element capable of binding SRY as set forth in Figure 1 at residues 4625-4634, a sequence corresponding to an upstream motif or cis element characteristic of GC2 as set forth in Figure 1 at residues 4678-4711, a sequence corresponding to an upstream motif or cis element capable of binding PEA3 as set forth in Figure 1 at residues 4765-4769, a sequence corresponding to an upstream motif or cis element capable of MIR as set forth in Figure 1 at residues 4759-4954, a sequence corresponding to an upstream motif or cis element capable of binding NF-HNF-1 as set forth in Figure 1 at residues 4923-4941, a sequence corresponding to a thyroid receptor upstream motif or cis element as set forth in Figure 1 at residues 5151-5156, and a sequence corresponding to an upstream motif or cis element capable of binding NFkB as set forth in Figure 1 at residues 5166-5175 because complete complementarity is not needed for stable hybridization. The TIGR nucleic acid molecules of the present invention therefore also include molecules that, although capable of specifically hybridizing with TIGR nucleic acid molecules may lack "complete complementarity."

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Any of a variety of methods may be used to obtain the above-described nucleic acid molecules (Elles, Methods in Molecular Medicine: Molecular Diagnosis of Genetic Diseases, Humana Press (1996), herein incorporated by reference). SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7, SEQ ID NO: 8, SEQ ID NO: 9, SEQ ID NO: 10, SEQ ID NO: 11, SEQ ID NO: 12, SEQ ID NO: 13, SEQ ID NO: 14, SEQ ID NO: 15, SEQ ID NO: 16, SEQ ID NO: 17, SEQ ID NO: 18, SEQ ID NO: 19, SEQ ID NO: 20, SEQ ID NO: 21, SEQ ID NO: 22, SEQ ID NO: 23, SEQ ID NO: 24, SEQ ID NO: 25,

SEQ ID NO: 33, sequences corresponding to an upstream motif or cis element characteristic of PRL-FP111 as set forth in Figure 1 at residues 370-388, and 4491-4502, respectively, a sequence corresponding to an upstream motif or cis element capable of binding GR/PR as set forth in Figure 1 at residues 433-445, sequences corresponding to an upstream shear stress motif or cis element as set forth in Figure 1 at residues 446-451, 1288-1293, 3597-3602, 4771-4776, and 5240-5245, respectively, sequences corresponding to glucocorticoid response upstream motif or cis element as set forth in Figure 1 at residues 574-600, 1042-1056, 2444-2468, 2442-2269, 3536-3563, 4574-4593, 4595-4614, 4851-4865, 4844-4864, 5079-5084, 5083-5111, respectively, a sequence corresponding to an upstream motif or cis element capable of binding CBE as set forth in Figure 1 at residues 735-746, a sequence corresponding to an upstream motif or cis element capable of binding NFE as set forth in Figure 1 at residues 774-795, a sequence corresponding to an upstream motif or cis element capable of binding KTF.1-CS as set forth in Figure 1 at residues 843-854, a sequence corresponding to an upstream motif or cis element capable of binding PRE is set forth in Figure 1 at residues 987-1026, a sequence corresponding to an upstream motif or cis element capable of binding ETF-EGFR as set forth in Figure 1 at residues 1373-1388, a sequence corresponding to an upstream motif or cis element capable of binding SRE-cFos as set forth in Figure 1 at residues 1447-1456, a sequence corresponding to an upstream motif or cis element capable of binding Alu as set forth in Figure 1 at residues 1331-1550, a sequence corresponding to an upstream motif or cis element capable of binding VBP as set forth in Figure 1 at residues 1786-1797, a sequence corresponding to an upstream motif or cis element capable of binding Malt-CS as set forth in Figure 1 at residues 1832-1841, sequences corresponding to an upstream motif or cis element capable of binding ERE as set forth in Figure 1 at residues 2167-2195, 3413-3429, and 3892-3896, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-mutagen as set forth in Figure 1 at residues 2329-2338, a sequence corresponding to an upstream motif or cis element capable of binding myc-PRF as set forth in Figure 1 at residues 2403-2416, sequences corresponding to an upstream motif or cis element capable of binding AP2 as set forth in Figure 1 at residues 2520-2535 and 5170-5187, respectively, sequences corresponding to an upstream motif or cis element capable of binding HSTF as set forth in Figure 1 at residues 2622-2635, and 5105-5132, respectively, a sequence corresponding to an upstream motif or cis element characteristic of SBF as set forth in Figure 1 at residues 2733-2743, sequences corresponding to an upstream motif or cis element capable of binding NF-1 as set forth in Figure 1 at residues 2923-2938, 4144-4157, and 4887-4900, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-MHCIIA/B as set forth in Figure 1 at residues 2936-2944, a sequence corresponding to an upstream motif or cis element capable of binding PEA1 as set forth in Figure 1 at residues 3285-3298, a sequence corresponding to an upstream motif or cis element

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capable of binding ICS as set forth in Figure 1 at residues 3688-3699, a sequence corresponding to an upstream motif or cis element capable of binding ISGF2 as set forth in Figure 1 at residues 4170-4179, a sequence corresponding to an upstream motif or cis element capable of binding zinc as set forth in Figure 1 at residues 4285-4293, a sequence corresponding to an upstream motif or cis element characteristic of CAP/CRP-galO as set forth in Figure 1 at residues 4379-4404, sequences corresponding to an upstream motif or cis element capable of binding AP1 as set forth in Figure 1 at residues 4428-4434, and 4627-4639, respectively, a sequence corresponding to an upstream motif or cis element capable of binding SRY as set forth in Figure 1 at residues 4625-4634, a sequence corresponding to an upstream motif or cis element characteristic of GC2 as set forth in Figure 1 at residues 4678-4711, a sequence corresponding to an upstream motif or cis element capable of binding PEA3 as set forth in Figure 1 at residues 4765-4769, a sequence corresponding to an upstream motif or cis element capable of MIR as set forth in Figure 1 at residues 4759-4954, a sequence corresponding to an upstream motif or cis element capable of binding NF-HNF-1 as set forth in Figure 1 at residues 4923-4941, a sequence corresponding to a thyroid receptor upstream motif or cis element as set forth in Figure 1 at residues 5151-5156, and a sequence corresponding to an upstream motif or cis element capable of binding NFkB as set forth in Figure 1 at residues 5166-5175 may be used to synthesize all or any portion of the TIGR promoter or any of the TIGR upstream motifs or portions the TIGR cDNA (Zamechik et al., Proc. Natl. Acad. Sci. (U.S.A.) 83:4143 (1986); Goodchild et al., Proc. Natl. Acad. Sci. (U.S.A.) 85:5507 (1988); Wickstrom et al., Proc. Natl. Acad. Sci. (U.S.A.) 85:1028; Holt, J.T. et al., Molec. Cell. Biol. 8:963 (1988); Gerwirtz, A.M. et al., Science 242:1303 (1988); Anfossi, G., et al., Proc. Natl. Acad. Sci. (U.S.A.) 86:3379 (1989); Becker, D., et al., EMBO J. 8:3679 (1989); all of which references are incorporated herein by reference).

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Automated nucleic acid synthesizers may be employed for this purpose. In lieu of such synthesis, the disclosed SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7, SEQ ID NO: 8, SEQ ID NO: 9, SEQ ID NO: 10, SEQ ID NO: 11, SEQ ID NO: 12, SEQ ID NO: 13, SEQ ID NO: 14, SEQ ID NO: 15, SEQ ID NO: 16, SEQ ID NO: 17, SEQ ID NO: 18, SEQ ID NO: 19, SEQ ID NO: 20, SEQ ID NO: 21, SEQ ID NO: 22, SEQ ID NO: 23, SEQ ID NO: 24, SEQ ID NO: 25, SEQ ID NO: 33, sequences corresponding to an upstream motif or cis element characteristic of PRL-FP111 as set forth in Figure 1 at residues 370-388, and 4491-4502, respectively, a sequence corresponding to an upstream motif or cis element capable of binding GR/PR as set forth in Figure 1 at residues 433-445, sequences corresponding to an upstream shear stress motif or cis element as set forth in Figure 1 at residues 446-451, 1288-1293, 3597-3602, 4771-4776, and 5240-5245, respectively, sequences corresponding to glucocorticoid response upstream motif or cis element as set forth in Figure 1 at residues 574-600, 1042-1056, 2444-2468, 2442-2269, 3536-3563, 4574-4593, 4595-

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4614, 4851-4865, 4844-4864, 5079-5084, 5083-5111, respectively, a sequence corresponding to an upstream motif or cis element capable of binding CBE as set forth in Figure 1 at residues 735-746, a sequence corresponding to an upstream motif or cis element capable of binding NFE as set forth in Figure 1 at residues 774-795, a sequence corresponding to an upstream motif or cis element capable of binding KTF.1-CS as set forth in Figure 1 at residues 843-854, a sequence corresponding to an upstream motif or cis element capable of binding PRE is set forth in Figure 1 at residues 987-1026, a sequence corresponding to an upstream motif or cis element capable of binding ETF-EGFR as set forth in Figure 1 at residues 1373-1388, a sequence corresponding to an upstream motif or cis element capable of binding SRE-cFos as set forth in Figure 1 at residues 1447-1456, a sequence corresponding to an upstream motif or cis element capable of binding Alu as set forth in Figure 1 at residues 1331-1550, a sequence corresponding to an upstream motif or cis element capable of binding VBP as set forth in Figure 1 at residues 1786-1797, a sequence corresponding to an upstream motif or cis element capable of binding Malt-CS as set forth in Figure 1 at residues 1832-1841, sequences corresponding to an upstream motif or cis element capable of binding ERE as set forth in Figure 1 at residues 2167-2195, 3413-3429, and 3892-3896, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-mutagen as set forth in Figure 1 at residues 2329-2338, a sequence corresponding to an upstream motif or cis element capable of binding myc-PRF as set forth in Figure 1 at residues 2403-2416, sequences corresponding to an upstream motif or cis element capable of binding AP2 as set forth in Figure 1 at residues 2520-2535 and 5170-5187, respectively, sequences corresponding to an upstream motif or cis element capable of binding HSTF as set forth in Figure 1 at residues 2622-2635, and 5105-5132, respectively, a sequence corresponding to an upstream motif or cis element characteristic of SBF as set forth in Figure 1 at residues 2733-2743, sequences corresponding to an upstream motif or cis element capable of binding NF-1 as set forth in Figure 1 at residues 2923-2938, 4144-4157, and 4887-4900, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-MHCIIA/B as set forth in Figure 1 at residues 2936-2944, a sequence corresponding to an upstream motif or cis element capable of binding PEA1 as set forth in Figure 1 at residues 3285-3298, a sequence corresponding to an upstream motif or cis element capable of binding ICS as set forth in Figure 1 at residues 3688-3699, a sequence corresponding to an upstream motif or cis element capable of binding ISGF2 as set forth in Figure 1 at residues 4170-4179, a sequence corresponding to an upstream motif or cis element capable of binding zinc as set forth in Figure 1 at residues 4285-4293, a sequence corresponding to an upstream motif or cis element characteristic of CAP/CRP-galO as set forth in Figure 1 at residues 4379-4404, sequences corresponding to an upstream motif or cis element capable of binding AP1 as set forth in Figure 1 at residues 4428-4434, and 4627-4639, respectively, a sequence corresponding to an upstream

motif or *cis* element capable of binding SRY as set forth in Figure 1 at residues 4625-4634, a sequence corresponding to an upstream motif or *cis* element characteristic of GC2 as set forth in Figure 1 at residues 4678-4711, a sequence corresponding to an upstream motif or *cis* element capable of binding PEA3 as set forth in Figure 1 at residues 4765-4769, a sequence corresponding to an upstream motif or *cis* element capable of MIR as set forth in Figure 1 at residues 4759-4954, a sequence corresponding to an upstream motif or *cis* element capable of binding NF-HNF-1 as set forth in Figure 1 at residues 4923-4941, a sequence corresponding to a thyroid receptor upstream motif or *cis* element as set forth in Figure 1 at residues 5151-5156, and a sequence corresponding to an upstream motif or *cis* element capable of binding NFkB as set forth in Figure 1 at residues 5166-5175 may be used to define a pair of primers that can be used with the polymerase chain reaction (Mullis, K. *et al.*, Cold Spring Harbor Symp. Quant. Biol. 51:263-273 (1986); Erlich H. *et al.*, EP 50,424; EP 84,796, EP 258,017, EP 237,362; Mullis, K., EP 201,184; Mullis K. *et al.*, US 4,683,202; Erlich, H., US 4,582,788; and Saiki, R. *et al.*, US 4,683,194)) to amplify and obtain any desired TIGR gene DNA molecule or fragment.

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The TIGR promoter sequence(s) and TIGR flanking sequences can also be obtained by incubating oligonucleotide probes of TIGR oligonucleotides with members of genomic human libraries and recovering clones that hybridize to the probes. In a second embodiment, methods of "chromosome walking," or 3' or 5' RACE may be used (Frohman, M.A. et al., Proc. Natl. Acad. Sci. (U.S.A.) 85:8998-9002 (1988), herein incorporated by reference); Ohara, O. et al., Proc. Natl. Acad. Sci. (U.S.A.) 86:5673-5677 (1989), herein incorporated by reference) to obtain such sequences.

II. Uses of the Molecules of the Invention in the Diagnosis and Prognosis of Glaucoma and Related Diseases

A particularly desired use of the present invention relates to the diagnosis of glaucoma, POAG, pigmentary glaucoma, high tension glaucoma and low tension glaucoma and their related diseases. Another particularly desired use of the present invention relates to the prognosis of glaucoma, POAG, pigmentary glaucoma, high tension glaucoma and low tension glaucoma and their related diseases. As used herein the term "glaucoma" includes both primary glaucomas, secondary glaucomas, juvenile glaucomas, congenital glaucomas, and familial glaucomas, including, without limitation, pigmentary glaucoma, high tension glaucoma and low tension glaucoma and their related diseases. As indicated above, methods for diagnosing or prognosing glaucoma suffer from inaccuracy, or require multiple examinations. The molecules of the present invention may be used to define superior assays for glaucoma. Quite apart from such usage, the molecules of the present invention may be used to diagnosis or predict an individual's sensitivity to elevated intraocular pressure upon administration of steroids such as

glucocorticoids or corticosteroids, or anti-inflammatory steroids). Dexamethasone, cortisol and prednisolone are preferred steroids for this purpose. Medical conditions such as inflammatory and allergic disorders, as well as organ transplantation recipients, benefit from treatment with glucocorticoids. Certain individuals exhibit an increased IOP response to such steroids (i.e., "steroid sensitivity"), which is manifested by an undesired increase in intraocular pressure. The present invention may be employed to diagnosis or predict such sensitivity, as well as glaucoma and related diseases.

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In a first embodiment, the TIGR molecules of the present invention are used to determine whether an individual has a mutation affecting the level (i.e., the concentration of TIGR mRNA or protein in a sample, etc.) or pattern (i.e., the kinetics of expression, rate of decomposition, stability profile, etc.) of the TIGR expression (collectively, the "TIGR response" of a cell or bodily fluid) (for example, a mutation in the TIGR gene, or in a regulatory region(s) or other gene(s) that control or affect the expression of TIGR), and being predictive of individuals who would be predisposed to glaucoma (prognosis), related diseases, or steroid sensitivity. As used herein, the TIGR response manifested by a cell or bodily fluid is said to be "altered" if it differs from the TIGR response of cells or of bodily fluids of normal individuals. Such alteration may be manifested by either abnormally increased or abnormally diminished TIGR response. To determine whether a TIGR response is altered, the TIGR response manifested by the cell or bodily fluid of the patient is compared with that of a similar cell sample (or bodily fluid sample) of normal individuals. As will be appreciated, it is not necessary to re-determine the TIGR response of the cell sample (or bodily fluid sample) of normal individuals each time such a comparison is made; rather, the TIGR response of a particular individual may be compared with previously obtained values of normal individuals.

In one sub-embodiment, such an analysis is conducted by determining the presence and/or identity of polymorphism(s) in the TIGR gene or its flanking regions which are associated with glaucoma, or a predisposition (prognosis) to glaucoma, related diseases, or steroid sensitivity. As used herein, the term "TIGR flanking regions" refers to those regions which are located either upstream or downstream of the TIGR coding region.

Any of a variety of molecules can be used to identify such polymorphism(s). In one embodiment, SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7, SEQ ID NO: 8, SEQ ID NO: 9, SEQ ID NO: 10, SEQ ID NO: 11, SEQ ID NO: 12, SEQ ID NO: 13, SEQ ID NO: 14, SEQ ID NO: 15, SEQ ID NO: 16, SEQ ID NO: 17, SEQ ID NO: 18, SEQ ID NO: 19, SEQ ID NO: 20, SEQ ID NO: 21, SEQ ID NO: 22, SEQ ID NO: 23, SEQ ID NO: 24, SEQ ID NO: 25, SEQ ID NO: 33, sequences corresponding to an upstream motif or *cis* element characteristic of PRL-FP111 as set forth in Figure 1 at residues 370-388, and 4491-4502, respectively, a sequence corresponding to an upstream motif or *cis*

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element capable of binding GR/PR as set forth in Figure 1 at residues 433-445, sequences corresponding to an upstream shear stress motif or cis element as set forth in Figure 1 at residues 446-451, 1288-1293, 3597-3602, 4771-4776, and 5240-5245, respectively, sequences corresponding to glucocorticoid response upstream motif or cis element as set forth in Figure 1 at residues 574-600, 1042-1056, 2444-2468, 2442-2269, 3536-3563, 4574-4593, 4595-4614, 4851-4865, 4844-4864, 5079-5084, 5083-5111, respectively, a sequence corresponding to an upstream motif or cis element capable of binding CBE as set forth in Figure 1 at residues 735-746, a sequence corresponding to an upstream motif or cis element capable of binding NFE as set forth in Figure 1 at residues 774-795, a sequence corresponding to an upstream motif or cis element capable of binding KTF.1-CS as set forth in Figure 1 at residues 843-854, a sequence corresponding to an upstream motif or cis element capable of binding PRE is set forth in Figure 1 at residues 987-1026, a sequence corresponding to an upstream motif or cis element capable of binding ETF-EGFR as set forth in Figure 1 at residues 1373-1388, a sequence corresponding to an upstream motif or cis element capable of binding SRE-cFos as set forth in Figure 1 at residues 1447-1456, a sequence corresponding to an upstream motif or cis element capable of binding Alu as set forth in Figure 1 at residues 1331-1550, a sequence corresponding to an upstream motif or cis element capable of binding VBP as set forth in Figure 1 at residues 1786-1797, a sequence corresponding to an upstream motif or cis element capable of binding Malt-CS as set forth in Figure 1 at residues 1832-1841, sequences corresponding to an upstream motif or cis element capable of binding ERE as set forth in Figure 1 at residues 2167-2195, 3413-3429, and 3892-3896, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-mutagen as set forth in Figure 1 at residues 2329-2338, a sequence corresponding to an upstream motif or cis element capable of binding myc-PRF as set forth in Figure 1 at residues 2403-2416, sequences corresponding to an upstream motif or cis element capable of binding AP2 as set forth in Figure 1 at residues 2520-2535 and 5170-5187, respectively, sequences corresponding to an upstream motif or cis element capable of binding HSTF as set forth in Figure 1 at residues 2622-2635, and 5105-5132, respectively, a sequence corresponding to an upstream motif or cis element characteristic of SBF as set forth in Figure 1 at residues 2733-2743, sequences corresponding to an upstream motif or cis element capable of binding NF-1 as set forth in Figure 1 at residues 2923-2938, 4144-4157, and 4887-4900, respectively, a sequence corresponding to an upstream motif or cis element capable of binding NF-MHCIIA/B as set forth in Figure 1 at residues 2936-2944, a sequence corresponding to an upstream motif or cis element capable of binding PEA1 as set forth in Figure 1 at residues 3285-3298, a sequence corresponding to an upstream motif or cis element capable of binding ICS as set forth in Figure 1 at residues 3688-3699, a sequence corresponding to an upstream motif or cis 35 element capable of binding ISGF2 as set forth in Figure 1 at residues 4170-4179, a sequence

corresponding to an upstream motif or cis element capable of binding zinc as set forth in Figure 1 at residues 4285-4293, a sequence corresponding to an upstream motif or cis element characteristic of CAP/CRP-galO as set forth in Figure 1 at residues 4379-4404, sequences corresponding to an upstream motif or cis element capable of binding AP1 as set forth in Figure 1 at residues 4428-4434, and 4627-4639, respectively, a sequence corresponding to an upstream motif or cis element capable of binding SRY as set forth in Figure 1 at residues 4625-4634, a sequence corresponding to an upstream motif or cis element characteristic of GC2 as set forth in Figure 1 at residues 4678-4711, a sequence corresponding to an upstream motif or cis element capable of binding PEA3 as set forth in Figure 1 at residues 4765-4769, a sequence corresponding to an upstream motif or cis element capable of MIR as set forth in Figure 1 at residues 4759-4954, a sequence corresponding to an upstream motif or cis element capable of binding NF-HNF-1 as set forth in Figure 1 at residues 4923-4941, a sequence corresponding to a thyroid receptor upstream motif or cis element as set forth in Figure 1 at residues 5151-5156, and a sequence corresponding to an upstream motif or cis element capable of binding NFkB as set forth in Figure 1 at residues 5166-5175 (or a sub-sequence thereof) may be employed as a marker nucleic acid molecule to identify such polymorphism(s).

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Alternatively, such polymorphisms can be detected through the use of a marker nucleic acid molecule or a marker protein that is genetically linked to (i.e., a polynucleotide that co-segregates with) such polymorphism(s). As stated above, the TIGR gene and/or a sequence or sequences that specifically hybridize to the TIGR gene have been mapped to chromosome 1q, 21-32, and more preferably to the TIGR gene located at chromosome 1, q21-27, and more preferably to the TIGR gene located at chromosome 1, q22-26, and most preferably to the TIGR gene located at chromosome 1, q24. In a preferred aspect of this embodiment, such marker nucleic acid molecules will have the nucleotide sequence of a polynucleotide that is closely genetically linked to such polymorphism(s) (e.g., markers located at chromosome 1, q19-25 (and more preferably chromosome 1, q23-25, and most preferably chromosome 1, q24.

Localization studies using a Stanford G3 radiation hybrid panel mapped the TIGR gene with the D1S2536 marker nucleic acid molecules at the D1S2536 locus with a LOD score of 6.0. Other marker nucleic acid molecules in this region include: D1S210; D1S1552; D1S2536; D1S2790; SHGC-12820; and D1S2558. Other polynucleotide markers that map to such locations are known and can be employed to identify such polymorphism(s).

The genomes of animals and plants naturally undergo spontaneous mutation in the course of their continuing evolution (Gusella, J.F., Ann. Rev. Biochem. 55:831-854 (1986)). A "polymorphism" in the TIGR gene or its flanking regions is a variation or difference in the sequence of the TIGR gene or its flanking regions that arises in some of the members of a

species. The variant sequence and the "original" sequence co-exist in the species' population. In some instances, such co-existence is in stable or quasi-stable equilibrium.

A polymorphism is thus said to be "allelic," in that, due to the existence of the polymorphism, some members of a species may have the original sequence (i.e. the original "allele") whereas other members may have the variant sequence (i.e. the variant "allele"). In the simplest case, only one variant sequence may exist, and the polymorphism is thus said to be diallelic. In other cases, the species' population may contain multiple alleles, and the polymorphism is termed tri-allelic, etc. A single gene may have multiple different unrelated polymorphisms. For example, it may have a di-allelic polymorphism at one site, and a multiple allelic polymorphism at another site.

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The variation that defines the polymorphism may range from a single nucleotide variation to the insertion or deletion of extended regions within a gene. In some cases, the DNA sequence variations are in regions of the genome that are characterized by short tandem repeats (STRs) that include tandem di- or tri-nucleotide repeated motifs of nucleotides. Polymorphisms characterized by such tandem repeats are referred to as "variable number tandem repeat" ("VNTR") polymorphisms. VNTRs have been used in identity and paternity analysis (Weber, J.L., U.S. Patent 5,075,217; Armour, J.A.L. et al., FEBS Lett. 307:113-115 (1992); Jones, L. et al., Eur. J. Haematol. 39:144-147 (1987); Horn, G.T. et al., PCT Application WO91/14003; Jeffreys, A.J., European Patent Application 370,719; Jeffreys, A.J., U.S. Patent 5,175,082); Jeffreys. A.J. et al., Amer. J. Hum. Genet. 39:11-24 (1986); Jeffreys. A.J. et al., Nature 316:76-79 (1985); Gray, I.C. et al., Proc. R. Acad. Soc. Lond. 243:241-253 (1991); Moore, S.S. et al., Genomics 10:654-660 (1991); Jeffreys, A.J. et al., Anim. Genet. 18:1-15 (1987); Hillel, J. et al., Anim. Genet. 20:145-155 (1989); Hillel, J. et al., Genet. 124:783-789 (1990)).

In an alternative embodiment, such polymorphisms can be detected through the use of a marker nucleic acid molecule that is physically linked to such polymorphism(s). For this purpose, marker nucleic acid molecules comprising a nucleotide sequence of a polynucleotide located within 1 mb of the polymorphism(s), and more preferably within 100 kb of the polymorphism(s), and most preferably within 10 kb of the polymorphism(s) can be employed. Examples of such marker nucleic acids are set out in SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7, SEQ ID NO: 8, SEQ ID NO: 9, SEQ ID NO: 10, SEQ ID NO: 11, SEQ ID NO: 12, SEQ ID NO: 13, SEQ ID NO: 14, SEQ ID NO: 15, SEQ ID NO: 16, SEQ ID NO: 17, SEQ ID NO: 18, SEQ ID NO: 19, SEQ ID NO: 20, SEQ ID NO: 21, SEQ ID NO: 22, SEQ ID NO: 23, SEQ ID NO: 24, SEQ ID NO: 25.

In another embodiment a marker nucleic acid will be used that is capable of specifically detecting TIGRmt1, TIGRmt2, TIGRmt3, TIGRmt4, TIGRmt5, TIGRmt11, TIGRsv1, or a combination of these mutations. Methods to detect base(s) substitutions, base(s) deletions and

base(s) additions are known in the art (i.e. methods to genotype an individual). For example, "Genetic Bit Analysis ("GBA") method is disclosed by Goelet, P. et al., WO 92/15712, herein incorporated by reference, may be used for detecting the single nucleotide polymorphisms of the present invention. GBA is a method of polymorphic site interrogation in which the nucleotide sequence information surrounding the site of variation in a target DNA sequence is used to design an oligonucleotide primer that is complementary to the region immediately adjacent to, but not including, the variable nucleotide in the target DNA. The target DNA template is selected from the biological sample and hybridized to the interrogating primer. This primer is extended by a single labeled dideoxynucleotide using DNA polymerase in the presence of two, and preferably all four chain terminating nucleoside triphosphate precursors. Cohen, D. et al., (PCT Application WO91/02087) describes a related method of genotyping.

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Other primer-guided nucleotide incorporation procedures for assaying polymorphic sites in DNA have been described (Komher, J. S. et al., Nucl. Acids. Res. 17:7779-7784 (1989), herein incorporated by reference; Sokolov, B. P., Nucl. Acids Res. 18:3671 (1990), herein incorporated by reference; Syvänen, A.-C., et al., Genomics 8:684 - 692 (1990), herein incorporated by reference; Kuppuswamy, M.N. et al., Proc. Natl. Acad. Sci. (U.S.A.) 88:1143-1147 (1991), herein incorporated by reference; Prezant, T.R. et al., Hum. Mutat. 1:159-164 (1992), herein incorporated by reference; Ugozzoli, L. et al., GATA 9:107-112 (1992), herein incorporated by reference; Nyrén, P. et al., Anal. Biochem. 208:171-175 (1993), herein incorporated by reference).

The detection of polymorphic sites in a sample of DNA may be facilitated through the use of nucleic acid amplification methods. Such methods specifically increase the concentration of polynucleotides that span the polymorphic site, or include that site and sequences located either distal or proximal to it. Such amplified molecules can be readily detected by gel electrophoresis or other means.

Another preferred method of achieving such amplification employs the polymerase chain reaction ("PCR") (Mullis, K. et al., Cold Spring Harbor Symp. Quant. Biol. 51:263-273 (1986); Erlich H. et al., European Patent Appln. 50,424; European Patent Appln. 84,796, European Patent Application 258,017, European Patent Appln. 237,362; Mullis, K., European Patent Appln. 201,184; Mullis K. et al., U.S. Patent No. 4,683,202; Erlich, H., U.S. Patent No. 4,582,788; and Saiki, R. et al., U.S. Patent No. 4,683,194), using primer pairs that are capable of hybridizing to the proximal sequences that define a polymorphism in its double-stranded form.

In lieu of PCR, alternative methods, such as the "Ligase Chain Reaction" ("LCR") may be used (Barany, F., Proc. Natl. Acad. Sci. (U.S.A.) 88:189-193 (1991). LCR uses two pairs of oligonucleotide probes to exponentially amplify a specific target. The sequences of each pair of oligonucleotides is selected to permit the pair to hybridize to abutting sequences of the same

strand of the target. Such hybridization forms a substrate for a template-dependent ligase. As with PCR, the resulting products thus serve as a template in subsequent cycles and an exponential amplification of the desired sequence is obtained.

LCR can be performed with oligonucleotides having the proximal and distal sequences of the same strand of a polymorphic site. In one embodiment, either oligonucleotide will be designed to include the actual polymorphic site of the polymorphism. In such an embodiment, the reaction conditions are selected such that the oligonucleotides can be ligated together only if the target molecule either contains or lacks the specific nucleotide that is complementary to the polymorphic site present on the oligonucleotide. Alternatively, the oligonucleotides may be selected such that they do not include the polymorphic site (see, Segev, D., PCT Application WO 90/01069).

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The "Oligonucleotide Ligation Assay" ("OLA") may alternatively be employed (Landegren, U. et al., Science 241:1077-1080 (1988)). The OLA protocol uses two oligonucleotides which are designed to be capable of hybridizing to abutting sequences of a single strand of a target. OLA, like LCR, is particularly suited for the detection of point mutations. Unlike LCR, however, OLA results in "linear" rather than exponential amplification of the target sequence.

Nickerson, D.A. et al., have described a nucleic acid detection assay that combines attributes of PCR and OLA (Nickerson, D.A. et al., Proc. Natl. Acad. Sci. (U.S.A.) 87:8923-8927 (1990). In this method, PCR is used to achieve the exponential amplification of target DNA, which is then detected using OLA. In addition to requiring multiple, and separate, processing steps, one problem associated with such combinations is that they inherit all of the problems associated with PCR and OLA.

Schemes based on ligation of two (or more) oligonucleotides in the presence of nucleic acid having the sequence of the resulting "di-oligonucleotide", thereby amplifying the di-oligonucleotide, are also known (Wu, D.Y. et al., Genomics 4:560 (1989)), and may be readily adapted to the purposes of the present invention.

Other known nucleic acid amplification procedures, such as allele-specific oligomers, branched DNA technology, transcription-based amplification systems, or isothermal amplification methods may also be used to amplify and analyze such polymorphisms (Malek, L.T. et al., U.S. Patent 5,130,238; Davey, C. et al., European Patent Application 329,822; Schuster et al., U.S. Patent 5,169,766; Miller, H.I. et al., PCT appln. WO 89/06700; Kwoh, D. et al., Proc. Natl. Acad. Sci. (U.S.A.) 86:1173 (1989); Gingeras, T.R. et al., PCT application WO 88/10315; Walker, G.T. et al., Proc. Natl. Acad. Sci. (U.S.A.) 89:392-396 (1992)). All the foregoing nucleic acid amplification methods could be used to predict or diagnose glaucoma.

The identification of a polymorphism in the TIGR gene, or flanking sequences up to about 5,000 base from either end of the coding region, can be determined in a variety of ways. By correlating the presence or absence of glaucoma in an individual with the presence or absence of a polymorphism in the TIGR gene or its flanking regions, it is possible to diagnose the predisposition (prognosis) of an asymptomatic patient to glaucoma, related diseases, or steroid sensitivity. If a polymorphism creates or destroys a restriction endonuclease cleavage site, or if it results in the loss or insertion of DNA (e.g., a VNTR polymorphism), it will alter the size or profile of the DNA fragments that are generated by digestion with that restriction endonuclease. As such, individuals that possess a variant sequence can be distinguished from those having the original sequence by restriction fragment analysis. Polymorphisms that can be identified in this manner are termed "restriction fragment length polymorphisms" ("RFLPs"). RFLPs have been widely used in human and animal genetic analyses (Glassberg, J., UK patent Application 2135774; Skolnick, M.H. et al., Cytogen. Cell Genet. 32:58-67 (1982); Botstein, D. et al., Ann. J. Hum. Genet. 32:314-331 (1980); Fischer, S.G et al. (PCT Application WO90/13668); Uhlen, M., PCT Application WO90/11369)). The role of TIGR in glaucoma pathogenesis indicates that the presence of genetic alterations (e.g., DNA polymorphisms) that affect the TIGR response can be employed to predict glaucoma.

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A preferred method of achieving such identification employs the single-strand conformational polymorphism (SSCP) approach. The SSCP technique is a method capable of identifying most sequence variations in a single strand of DNA, typically between 150 and 250 nucleotides in length (Elles, Methods in Molecular Medicine: Molecular Diagnosis of Genetic Diseases, Humana Press (1996), herein incorporated by reference); Orita et al., Genomics 5: 874-879 (1989), herein incorporated by reference). Under denaturing conditions a single strand of DNA will adopt a conformation that is uniquely dependent on its sequence conformation. This conformation usually will be different, even if only a single base is changed. Most conformations have been reported to alter the physical configuration or size sufficiently to be detectable by electrophoresis. A number of protocols have been described for SSCP including, but not limited to Lee et al., Anal. Biochem. 205: 289-293 (1992), herein incorporated by reference; Suzuki et al., Anal. Biochem. 192: 82-84 (1991), herein incorporated by reference; Lo et al., Nucleic Acids Research 20: 1005-1009 (1992), herein incorporated by reference; Sarkar et al., Genomics 13: 441-443 (1992), herein incorporated by reference).

In accordance with this embodiment of the invention, a sample DNA is obtained from a patient. In a preferred embodiment, the DNA sample is obtained from the patient's blood. However, any source of DNA may be used. The DNA is subjected to restriction endonuclease digestion. TIGR is used as a probe in accordance with the above-described RFLP methods. By comparing the RFLP pattern of the TIGR gene obtained from normal and glaucomatous patients,

one can determine a patient's predisposition (prognosis) to glaucoma. The polymorphism obtained in this approach can then be cloned to identify the mutation at the coding region which alters the protein's structure or regulatory region of the gene which affects its expression level. Changes involving promoter interactions with other regulatory proteins can be identified by, for example, gel shift assays using HTM cell extracts, fluid from the anterior chamber of the eye, serum, etc. Interactions of TIGR protein in glaucomatous cell extracts, fluid from the anterior chamber of the eye, serum, etc. can be compared to control samples to thereby identify changes in those properties of TIGR that relate to the pathogenesis of glaucoma. Similarly such extracts and fluids as well as others (blood, etc.) can be used to diagnosis or predict steroid sensitivity.

Several different classes of polymorphisms may be identified through such methods. Examples of such classes include: (1) polymorphisms present in the TIGR cDNA of different individuals; (2) polymorphisms in non-translated TIGR gene sequences, including the promoter or other regulatory regions of the TIGR gene; (3) polymorphisms in genes whose products interact with TIGR regulatory sequences; (4) polymorphisms in gene sequences whose products interact with the TIGR protein, or to which the TIGR protein binds.

In an alternate sub-embodiment, the evaluation is conducted using oligonucleotide "probes" whose sequence is complementary to that of a portion of SEQ ID NO: 1, SEQ ID NO: 2 SEQ ID NO: 3, SEQ ID NO: 4, or SEQ ID NO: 5. Such molecules are then incubated with cell extracts of a patient under conditions sufficient to permit nucleic acid hybridization.

In one sub-embodiment of this aspect of the present invention, one can diagnose or predict glaucoma, related diseases and steroid sensitivity by ascertaining the TIGR response in a biopsy (or a macrophage or other blood cell sample), or other cell sample, or more preferably, in a sample of bodily fluid (especially, blood, serum, plasma, tears, buccal cavity, etc.). Since the TIGR gene is induced in response to the presence of glucocorticoids, a highly preferred embodiment of this method comprises ascertaining such TIGR response prior to, during and/or subsequent to, the administration of a glucocorticoid. Thus, by way of illustration, glaucoma could be diagnosed or predicted by determining whether the administration of a glucocorticoid (administered topically, intraocularly, intramuscularly, systemically, or otherwise) alters the TIGR response of a particular individual, relative to that of normal individuals. Most preferably, for this purpose, at least a "TIGR gene-inducing amount" of the glucocorticoid will be provided. As used herein, a TIGR gene-inducing amount of a glucocorticoid is an amount of glucocorticoid sufficient to cause a detectable induction of TIGR expression in cells of glaucomatous or non-glaucomatous individuals.

Generating Cells, Vectors, and Expressed Proteins Using Agents of the Invention

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The present invention also relates to methods for obtaining a recombinant host cell, especially a mammalian host cell, comprising introducing into a host cell exogenous genetic material comprising a nucleic acid of the invention. The present invention also relates to an insect cell comprising a recombinant vector having a nucleic acid of the invention. The present invention also relates to methods for obtaining a recombinant host cell, comprising introducing exogenous genetic material comprising a nucleic acid of the invention via homologous recombination. Through homologous recombination, the promoter and 5' flanking sequences of the TIGR gene described here can be used in gene activation methods to produce a desired gene product in host cells (see, for example, U.S. Patent 5,733,746, specifically incorporated herein by reference). The specific expression of the TIGR gene in TM cells afforded by the TIGR promoter region DNA can, thus, be transferred via homologous recombination to express other gene products in a similar fashion. Some of these other gene products may be therapeutic proteins that address diseases related to increased IOP or glaucoma. Methods for selecting and using the promoter and 5' flanking sequence for the gene targeting technique involved in the gene activation method are known in the art. Depending upon the nature of the modification and associated targeting construct, various techniques may be employed for identifying targeted integration. Conveniently, the DNA may be digested with one or more restriction enzymes and the fragments probed with an appropriate DNA fragment, which will identify the properly sized restriction fragment associated with integration.

The sequence to be integrated into the host may be introduced by any convenient means, which includes calcium precipitated DNA, spheroplast fusion, transformation, electroporation, biolistics, lipofection, microinjection, or other convenient means. Where an amplifiable gene is being employed, the amplifiable gene may serve as the selection marker for selecting hosts into which the amplifiable gene has been introduced. Alternatively, one may include with the amplifiable gene another marker, such as a drug resistance marker, e.g. neomycin resistance (G418 in mammalian cells), hygromycin resistance etc., or an auxotrophy marker (HIS3, TRP1, LEU2, URA3, ADE2, LYS2, etc.) for use in yeast cells.

For example, homologous recombination constructs can be prepared where the amplifiable gene will be flanked, normally on both sides, with DNA homologous with the DNA of the target region, here the TIGR sequences. Depending upon the nature of the integrating DNA and the purpose of the integration, the homologous DNA will generally be within 100 kb, usually 50 kb, preferably about 25 kb, of the transcribed region of the target gene, more preferably within 2 kb of the target gene. The homologous DNA may include the 5'-upstream region outside of the transcriptional regulatory region or enhancer sequences, transcriptional

initiation sequences, adjacent sequences, or the like. The homologous region may include a portion of the coding region, where the coding region may be comprised only of an open reading frame or of combination of exons and introns. The homologous region may also comprise all or a portion of an intron, where all or a portion of one or more exons may also be present. Alternatively, the homologous region may comprise the 3'-region, so as to comprise all or a portion of the transcriptional termination region, or the region 3' of this region. The homologous regions may extend over all or a portion of the target gene or be outside the target gene comprising all or a portion of the transcriptional regulatory regions and/or the structural gene.

The integrating constructs may be prepared in accordance with conventional ways, where sequences may be synthesized, isolated from natural sources, manipulated, cloned, ligated, subjected to in vitro mutagenesis, primer repair, or the like. At various stages, the joined sequences may be cloned, and analyzed by restriction analysis, sequencing, or the like. Usually during the preparation of a construct where various fragments are joined, the fragments, intermediate constructs and constructs will be carried on a cloning vector comprising a replication system functional in a prokaryotic host, e.g., *E. coli*, and a marker for selection, e.g., biocide resistance, complementation to an auxotrophic host, etc. Other functional sequences may also be present, such as polylinkers, for ease of introduction and excision of the construct or portions thereof, or the like. A large number of cloning vectors are available such as pBR322, the pUC series, etc. These constructs may then be used for integration into the primary host.

DNA comprising a nucleic acid of the invention can be introduced into a host cell by a variety of techniques that include calcium phosphate/DNA co-precipitates, microinjection of DNA into the nucleus, electroporation, yeast protoplast fusion with intact cells, transfection, polycations, e.g., polybrene, polyornithine, etc., or the like. The DNA may be single or double stranded DNA, linear or circular. The various techniques for transforming cells are well known (see Keown et al., Methods Enzymol. (1989), Keown et al., Methods Enzymol. 185:527-537 (1990); Mansour et al., Nature 336:348-352, (1988); all of which are herein incorporated by reference in their entirety).

In a preferred aspect, the invention relates to recombinant insect vectors and insect cells comprising a nucleic acid of the invention. In a particularly preferred aspect, a Baculovirus expression vector is used, introduced into an insect cell, and recombinant TIGR protein expressed. The recombinant TIGR protein may be the full length protein from human TM endothelial cells, a fusion protein comprising a substantial fragment of the full length protein, for example, at least about 20 contiguous amino acids to about 100 contiguous amino acids of the full length protein, or a variant TIGR protein or fusion protein produced by site-directed mutagenesis, DNA shuffling, or a similar technique. Generally, the variant TIGR proteins and the fusion proteins will retain at least one structural or functional characteristic of the full length

TIGR protein, such as the ability to bind the same antibody, the presence of the substantially similar leucine zipper region, or the ability to bind the same ligand or receptor on TM cells (see Nguyen et al., J. Biol. Chem. 273:6341-6350 (1998), specifically incorporated herein by reference). Nucleic acids comprising the leucine zipper-encoding regions of the TIGR gene can be identified by methods known in the art and can be used in combination with recombinant or synthetic methods to create ligand-receptor assays.

Examples of the preferred, recombinant insect vector, host cell, and TIGR protein of the invention were generated by ligating TIGR cDNA into the PVL1393 vector [Invitrogen]. This vector was transferred into Sf9 cells, the TIGR protein expressed and then purified (see U.S. Patent 5,789,169 and Nguyen *et al.*, *J. Biol. Chem.* 273:6341-6350 (1998), both of which are specifically incorporated herein by reference in their entirety). An SDS-PAGE gel of the resulting proteins showed protein bands in the 55 kDa range, which were sequenced to confirm correct identity.

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In preferred embodiments of the vectors, cells and related methods of the invention, a TIGR fusion protein with GFP (green fluorescent protein) can be expressed in a TM cell line (see Nguyen, et al., J. Biol. Chem. 273:6341-6350 (1998) and the references cited therein for primary TM cell culture and transfection methods). Transformed, cultured TM cells at log phase were transfected with a TIGR-GFP fusion protein-encoding vector. The vector includes the CMV promoter to allow high expression, TIGR cDNA from the first ATG to the end of the proteinencoding region, a fluorescent protein tag (GFP) fused to the carboxy terminus of the TIGRencoding sequence, and the G418 resistance gene. These elements, and their use, is known in the art or provided by this disclosure and its incorporated references. The construct is termed TIGR1-GFP. The transfection was performed using calcium phosphate or Lipofectin techniques, as known in the art. Incubation at growth condition of 37°C, 8% CO₂, for 6-18 hours followed. After the transfection, the DNA media was replaced by fresh growth media including G418, which was changed twice weekly, until resistant colonies of cells outgrew the monolayer cells (about 10-15 days). The cell colonies were collected and propagated several passes to select for resistant, transformed cells. The expression of fluorescent TIGR-GFP fusion protein was tested for after several passes. One out of twenty selected colonies expressed high levels of the TIGR-GFP fusion protein.

In other preferred embodiments of the cells and methods of the invention, a transformed, immortalized TM cell line can be prepared using an SV40-derived vector. Primary cultured TM cells are transfected with an SV40 vector with a defect in the PsvOri, as known in the art. Briefly, primary cultured cells at log phase are transfected with PsvOri DNA using calcium phosphate or Lipofectin and incubated at growth condition of 37°C, 8% CO₂ for 6-18 hours. The DNA media was replaced by fresh growth media and changed twice weekly until colonies of

immortalized cells outgrow the dying monolayer (about 10-15 days). The cell colonies are collected and propagated several passes to select for transformed cells.

III. Methods of Administration

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Some of the agents of the present invention can be formulated according to known methods to prepare pharmacologically acceptable compositions, whereby these materials, or their functional derivatives, having the desired degree of purity are combined in admixture with a physiologically acceptable carrier, excipient, or stabilizer. Such materials are non-toxic to recipients at the dosages and concentrations employed. The active component of such compositions may be agents, analogs or mimetics of such molecules. Where nucleic acid molecules are employed, such molecules may be sense, antisense or triplex oligonucleotides of the TIGR promoter, TIGR cDNA, TIGR intron, TIGR exon or TIGR gene.

A composition is said to be "pharmacologically acceptable" if its administration can be tolerated by a recipient patient. An agent is physiologically significant if its presence results in a detectable change in the physiology of a recipient patient.

Suitable vehicles and their formulation, inclusive of other human proteins, e.g., human serum albumin, are described, for example, in Remington's Pharmaceutical Sciences (16th ed., Osol, A., Ed., Mack, Easton PA (1980)).

If the composition is to be water soluble, it may be formulated in a buffer such as phosphate or other organic acid salt preferably at a pH of about 7 to 8. If the composition is only partially soluble in water, it may be prepared as a microemulsion by formulating it with a nonionic surfactant such as Tween, Pluronics, or PEG, e.g., Tween 80, in an amount of, for example, 0.04-0.05% (w/v), to increase its solubility. The term "water soluble" as applied to the polysaccharides and polyethylene glycols is meant to include colloidal solutions and dispersions. In general, the solubility of the cellulose derivatives is determined by the degree of substitution of ether groups, and the stabilizing derivatives useful herein should have a sufficient quantity of such ether groups per anhydroglucose unit in the cellulose chain to render the derivatives water soluble. A degree of ether substitution of at least 0.35 ether groups per anhydroglucose unit is generally sufficient. Additionally, the cellulose derivatives may be in the form of alkali metal salts, for example, the Li, Na, K or Cs salts.

Optionally other ingredients may be added such as antioxidants, e.g., ascorbic acid; low molecular weight (less than about ten residues) polypeptides, e.g., polyarginine or tripeptides; proteins, such as serum albumin, gelatin, or immunoglobulins; hydrophilic polymers such as polyvinyl pyrrolidone; amino acids, such as glycine, glutamic acid, aspartic acid, or arginine;

monosaccharides, disaccharides, and other carbohydrates including cellulose or its derivatives, glucose, mannose, or dextrins; chelating agents such as EDTA; and sugar alcohols such as mannitol or sorbitol.

Additional pharmaceutical methods may be employed to control the duration of action. Controlled or sustained release preparations may be achieved through the use of polymers to complex or absorb the TIGR molecule(s) of the composition. The controlled delivery may be exercised by selecting appropriate macromolecules (for example polyesters, polyamino acids, polyvinyl pyrrolidone, ethylenevinylacetate, methylcellulose, carboxymethylcellulose, or protamine sulfate) and the concentration of macromolecules as well as the methods of incorporation in order to control release.

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Sustained release formulations may also be prepared, and include the formation of microcapsular particles and implantable articles. For preparing sustained-release compositions, the TIGR molecule(s) of the composition is preferably incorporated into a biodegradable matrix or microcapsule. A suitable material for this purpose is a polylactide, although other polymers of poly-(a-hydroxycarboxylic acids), such as poly-D-(-)-3-hydroxybutyric acid (EP 133,988A), can be used. Other biodegradable polymers include poly(lactones), poly(orthoesters), polyamino acids, hydrogels, or poly(orthocarbonates) poly(acetals). The polymeric material may also comprise polyesters, poly(lactic acid) or ethylene vinylacetate copolymers. For examples of sustained release compositions, see U.S. Patent No. 3,773,919, EP 58,481A, U.S. Patent No. 3,887,699, EP 158,277A, Canadian Patent No. 1176565, Sidman, U. et al., Biopolymers 22:547 (1983), and Langer, R. et al., Chem. Tech. 12:98 (1982).

Alternatively, instead of incorporating the TIGR molecule(s) of the composition into polymeric particles, it is possible to entrap these materials in microcapsules prepared, for example, by coacervation techniques or by interfacial polymerization, for example, hydroxymethylcellulose or gelatine-microcapsules and poly(methylmethacylate) microcapsules, respectively, or in colloidal drug delivery systems, for example, liposomes, albumin microspheres, microemulsions, nanoparticles, and nanocapsules or in macroemulsions. Such techniques are disclosed in Remington's Pharmaceutical Sciences (1980).

In an alternative embodiment, liposome formulations and methods that permit intracellular uptake of the molecule will be employed. Suitable methods are known in the art, see, for example, Chicz, R.M. et al. (PCT Application WO 94/04557), Jaysena, S.D. et al. (PCT Application WO93/12234), Yarosh, D.B. (U.S. Patent No. 5,190,762), Callahan, M.V. et al. (U.S. Patent No. 5,270,052) and Gonzalezro, R.J. (PCT Application 91/05771), all herein incorporated by reference.

PCT/US00/00559 WO 00/42220

Having now generally described the invention, the same will be more readily understood through reference to the following examples which are provided by way of illustration, and are not intended to be limiting of the present invention, unless specified.

EXAMPLE 1

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Illustrative Single Strand Conformational Polymorphism Assay

Single strand conformational polymorphism (SSCP) screening is carried out according to the procedure of Hue et al., The Journal of Investigative Ophthalmology 105.4: 529-632 (1995), herein incorporated by reference. SSCP primers are constructed corresponding to sequences found within the TIGR promoter and two of exons of TIGR. The following primers are constructed: forward primer "Sk-1a": 5'-TGA GGC TTC CTC TGG AAA C-3' (SEQ ID NO: 6); reverse primer "ca2": 5'-TGA AAT CAG CAC ACC AGT AG-3' (SEQ ID NO: 7); forward primer "CA2": 5'-GCA CCC ATA CCC CAA TAA TAG-3' (SEQ ID NO: 8); reverse primer "Pr+1": 5'-AGA GTT CCC CAG ATT TCA CC-3' (SEQ ID NO: 9); forward primer "Pr-1": 5'-ATC TGG GGA ACT CTC AG-3' (SEQ ID NO: 10); reverse primer "Pr+2(4A2)": 5'-TAC AGT TGT TGC AGA TAC G-3' (SEQ ID NO: 11); forward primer "Pr-2(4A)": 5'-ACA ACG TAT CTG CAA CAA CTG-3' (SEQ ID NO: 12); reverse primer "Pr+3(4A)": 5'-TCA GGC TTA ACT GCA GAA CC-3' (SEQ ID NO: 13); forward primer "Pr-3(4A)": 5'-TTG GTT CTG CAG TTA AGC C-3' (SEQ ID NO: 14); reverse primer "Pr+2(4A1)": 5'-AGC AGC ACA AGG GCA ATC C-3' (SEQ ID NO: 15); reverse primer "Pr+1(4A)": 5'-ACA GGG CTA TAT TGT 20 GGG-3' (SEQ ID NO: 16); forward primer "KS1X": 5'-CCT GAG ATG CCA GCT GTC C-3' (SEQ ID NO: 17); reverse primer "SK1XX": 5'-CTG AAG CAT TAG AAG CCA AC-3' (SEQ ID NO: 18); forward primer "KS2a1": 5'-ACC TTG GAC CAG GCT GCC AG-3' (SEQ ID NO: 19); reverse primer "SK3" 5'-AGG TTT GTT CGA GTT CCA G-3' (SEQ ID NO: 20); forward primer "KS4": 5'-ACA ATT ACT GGC AAG TAT GG-3' (SEQ ID NO: 21); reverse primer 25 "SK6A": 5'-CCT TCT CAG CCT TGC TAC C-3' (SEQ ID NO: 22); forward primer "KS5": 5'-ACA CCT CAG CAG ATG CTA CC-3' (SEQ ID NO: 23); reverse primer "SK8": 5'-ATG GAT GAC TGA CAT GGC C-3' (SEQ ID NO: 24); forward primer "KS6": 5'-AAG GAT GAA CAT GGT CAC C-3' (SEQ ID NO: 25).

The locations of primers: Sk-1a, ca2, CA2, Pr+1, Pr-1, Pr+2(4A2), Pr-2(4A), Pr+3(4A), Pr-3 (4A), Pr-3(4A), Pr+2(4A1), and Pr+1(4A) are diagramatically set forth in Figure 4. The location of primers: KS1X, SK1XX, Ks2a1, SK3, KS4, SK6A, KS5, SK8, and KS6 are diagramatically set forth in Figure 5.

Families with a history of POAG in Klamath Falls, Oregon, are screened by SSCP according to the method of Hue et al., The Journal of Investigative Ophthalmology 105.4: 529-

632 (1995), herein incorporated by reference). SSCP primers SK-1a, ca2, CA2, Pr+1, Pr-2(4A), Pr+3(4A), SK1XX, and KS6 detect single strand conformational polymorphisms in this population. An SSCP is detected using SSCP primers Pr+3(4A) and Pr-2(4A). 70 family members of the Klamath Fall, Oregon are screened with these primers and the results are set forth in Table 1.

TABLE 1

		Total	SSCP+	SSCP-
	Glaucoma positive individuals 1	12	12	0
10	Glaucoma negative individuals	13	. 0	13
	Spouses (glaucoma negative)	16	2	14
	Others ²	29	6	23

^{1 =} glaucoma positive individuals as determined by IOP of greater than 25 mmHg

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A second SSCP is detected using SSCP primers Pr+1 and CA2. 14 family members of the Klamath Fall, Oregon are screened with these primers. A characteristic polymorphism is found in the 6 affected family members but absent in the 8 unaffected members. A third SSCP is detected using SSCP primers ca2 and sk-1a. The same 14 family members of the Klamath Fall, Oregon that are screened with Pr+1 and CA2 are screened with ca2 and sk-1a primers. A characteristic polymorphism is found in the 6 affected family members but absent in the 8 unaffected members. A fourth SSCP is detected using SSCP primers KS6 and SK1XX. 22 family members of the Klamath Fall, Oregon and 10 members of a Portland, Oregon pedigree are screened with these primers. A polymorphism is found in exon 3. The results are as set forth in Table 2.

TABLE 2

	Total	SSCP+	SSCP-
Klamath Fall, Oregon Glaucoma positive individuals 1	3	3	0
Glaucoma negative individuals	6	0	6
Others ²	13	6	7
Portland, Oregon Glaucoma positive individuals 1	6	6	. 0
35 Glaucoma negative individuals	4	0	4

^{2 =} unidentified glaucoma due to the age of the individual.

Others ² 0 0 0

1 = glaucoma positive individuals as determined by IOP of greater than 25 mmHg

2 = unidentified glaucoma due to the age of the individual.

EXAMPLE 2

TIGR Homologies

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A novel "myosin-like" acidic protein termed myocilin is expressed predominantly in the photoreceptor cells of retina and is localized particularly in the rootlet and basal body of connecting cilium (Kubota *et al.*, Genomics 41: 360-369 (1997), herein incorporated by reference). The myocilin gene is mapped to human chromosome Iq23-q24. The coding region of myocilin is 100 percent homologous with TIGR.

Homology searches are performed by GCG (Genetics Computer Group, Madison, WI) and include the GenBank, EMBL, Swiss-Prot databases and EST analysis. Using the Blast search, the best fits are found with a stretch of 177 amino acids in the carboxy terminals for an extracellular mucus protein of the olfactory, olfactomedin and three olfactomedin-like species. The alignment presented in Figure 6 shows the TIGR homology (SEQ ID NO. 27) to an expressed sequence tag (EST) sequence from human brain (ym08h12.r1)(SEQ ID NO. 28)(The WashU-Merck EST Project, 1995); the Z domain of olfactomedin-related glycoprotein from rat brain (1B426bAMZ)(SEQ ID NO. 29)(Danielson et al., Journal of Neuroscience Research 38: 468-478 (1994), herein incorporated by reference) and the olfactomedin from olfactory tissue of bullfrogs (ranofm) (SEQ ID NO. 30)(Yokoe and Anholt, Proc. Natl. Acad. Sci. 90: 4655-4659 (1993), herein incorporated by reference; Snyder and Anholt, Biochemistry 30: 9143-9153 (1991), herein incorporated by reference). These domains share very similar amino acid positions as depicted in the consensus homology of Figure 6 (SEQ ID NO. 31), with the exception being the truncated human clone in which the position with respect to its full length sequence has not been established. No significant homology is found for the amino termini of these molecules.

EXAMPLE 3

Identification of TIGRmt11

DNA samples were obtained from individuals noted for having elevated IOP in response to the administration of topical corticosteroids. Typically, the "Armaly" criteria is used to register IOP changes.

Genomic DNA from blood or buccal swabs were used for PCR amplification. The PCR reaction includes 95° C for 30 sec, for denaturation, 55° C for 30 sec, for annealing and 72° C for

30 sec for synthosis. The reaction was performed for 30 cycles with an additional cycle of 72° C for 5 min at the end.

The primer pair for the PCR reaction can include any pair that amplifies a specific region targeted for analyzing mutants or polymorphisms. Preferably, the amplified region will be from about 500 base pairs 5' of the start of transcription to the start of translation. More preferably, it will include an amplified region about 200 bp 5' of the start of transcription to about 10 base pairs 5' to the start of translation. Methods for determining amplification primer sequences from within a known sequence region are well known in the art. Examplary methods include, but are not limited to, computer generated searches using programs such as Primer3 (www-genome.wi.mit.edu/cgi-bin/primer/primer3.cgi), STSPipeline (www-genome.wi.mit.edu/cgi-bin/www-STS_Pipeline), or GeneUp (Pesole, et al., BioTechniques 25:112-123 (1998)).

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In an especially preferred embodiment, this amplified region will be from position 5044 of SEQ ID NO: 3 to about 5327 of SEQ ID NO: 3, which will thus employ primers of the sequence of about 5044 to about 5066 and the sequence of about 5309 to about 5327 of SEQ ID NO: 3, or the complement. In one embodiment, the complement of the sequence from about 5309 to about 5327 is used as one of the primers and the sequence from about 5044 to about 5066 is used as the other primer.

For this example, the following primers were used: forward primer CA-2R (SEQ ID NO: 35 – 5' AACTATTATT GGGGTATGGG) and reverse primer Sk-la (SEQ ID NO: 36 - 5' TTGGTGAGGC TTCCTCTGC). The primers were labeled with a fluorescent dye IRD-800 by Li-Cor Technology and the PCR product (about 300 bp) was denatured by heat and subject to BESS assays to detect mutations.

BESS, or Base Excision Sequence Scanning, employed specific restriction enzyme that cleaves T position of single strand DNA. The cleavage will produce DNA fragments that could be observed by acrylamide gels. Based on this, a "T mutation" will produce different cleavage pattern for the mutated strand compared to the normal strand. Since 95% of mutations involve a T mutation, this method is very practical. In addition to BESS, the amplified fragments can also be sequenced or compared by hybridization methods (microarray hybridization techniques or the sequencing-by-hybridization technique) in order to determine the exact nucleotide sequence, as known in the art.

Using this assay, patients exhibiting an increased IOP in response to topical corticosteroid treatments had an elevated level of a T mutation in one particular position, at about

160 bases 5' to the start of the TIGR coding region. The presence of this particular mutation, called TIGRmt11, therefore, indicated a specific genetic linkage to steroid sentivity that manifests in atleast a higher risk of increased IOP, and thus glaucoma, in repsonse to steroid treatment.

TABLE 3

Subject	Duration of CS Treatment	IOP (OD/OS)	Genotype (mt.11)
1	1 year	38/30	+/-
2	3 weeks	25/28	+/+
3	2 weeks	28/28	+/+

CS= corticosteroid, topical treatment

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(1 year) CS treatment 38/30 mm Hg, OD/OS; (3 weeks) CS treatment 25/28 mm Hg, OD/OS; (2 weeks) CS treatment 28/38 mm Hg, OD/OS

The sequence in SEQ ID NO: 33 (CAAACAGACT TCCGGAAGGT) identifies bases immediately adjacent to the single base polymorphism, which represents bases 5101 to 5120 of SEQ ID NO: 1, except that the underlined C in the TIGRmt11 sequence variant is substituted for the 'wild type' T, found in SEQ ID NO: 1.

EXAMPLE 4

Verification of Linkage Between TIGRmt11 and Risk of Glaucoma

Subjects are given standard topical dexamethasone eye drops (0.1%) four times a day, for four weeks. Pre-treatment and post-treatment IOP readings are taken and patients are classified as having high (>16mmHg), intermediate (6-16mmHg) or low (<6mmHg) IOP responses under the "Armaly" criteria. DNA samples are obtained from four subjects having high or intermediate IOP changes. Samples from several non-responder patients were also taken. The DNA samples were analyzed for the presence of the TIGRmt11 variant sequence, as discussed above. The results are given in Table 4.

TABLE 4

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Subject	Age	Classification	CS-IOP Response	Genotype (mt.11)
1	47	OHT	Intermediate	+/+
2	28	POAG	High	+/+
3	46	POAG/OHT	High	+/+
4	15	Stevens-Johnson	High	+/+
5	Nr	Normal	Low	-/-
6	Nr	Normal	Low	- /-
7	Nr	Normal	Low	-/-

OHT = Ocular Hypertensive (began with a mild IOP elevation, no POAG)

POAG = Original diagnosis is primary open-angle glaucoma

POAG/OHT = Converted to POAG, from original diagnosis OHT

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The data obtained indicates the association of TIGRmt.11 and the response to topical CS. Clearly, all the subjects with clinically identifiable responses to the CS treatment possessed the TIGRmt11 variant sequence while none of the subjects with the 'wild type's equence, or a sequence that did not possess the TIGRmt11 variant, did not.

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While the invention has been described in connection with specific embodiments thereof, it will be understood that it is capable of further modifications and this application is intended to cover any variations, uses, or adaptations of the invention following, in general, the principles of the invention and including such departures from the present disclosure as come within known or customary practice within the art to which the invention pertains and as may be applied to the essential features herein before set forth and as follows in the scope of the appended claims.

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EXAMPLE 5

TIGR 5' Region /Promoter Activity Assays and Methods for Detecting Cell Components Binding
to TIGR Gene Sequences

The 283 base 5' fragment of the TIGR 5' region (SEQ ID NO: 37) is amplified from genomic DNA of a normal human subject as discusssed in Example 3. The DNA is cloned into a pSEAP vector (Clontech, CA) so that transcription was under the control of the TIGR sequences. More specifically, pSEAP2-enhancer vector is digested with Srfl to produce a blunt end. The 283 bp fragment is blunt end ligated to the vector using T4 ligase for 2 hours. The vector is transfected into DH5 cells to establish a plasmid clone, TMRE-1/pSEAP. PCR sequencing is used to verify the clone sequence as correct. Similarly, the equivalent genomic fragment amplified from a steroid responder sample, identified as mutant TIGR.mt11, is cloned in pSEAP2-enhancer and called clone TIGR.mt11/pSEAP.

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HTM cells, Cos cells (COS-7), and HeLa cells (HeLa 229) are seeded at 10⁶ cells per well and incubated with 1 ug of the plasmid DNA to transfect using Lipofectin reagent (GIBCO, LifeTechnologies, MD). Some samples of cells were also treated with dexamethasone (DEX). Then, 24, 48, and 72 hours after transfection, AP activity was measured. Cells are collected using a buffer containing L-homoarginine and the chemiluminescent substrate CSPD and a chemiluminescent enhancer added to the samples. The expression levels are then recorded using a luminometer or by brief exposure to X-ray film. The results show that AP activity is present in HTM transfected cells within 24 hours and that the levels increase over 50 fold 48 and 72 hours after transfection. In contrast, HeLa and Cos cells show insignificant AP activity at the same time points.

To detect cell components that specifically bind to TIGR sequences, the amplified genomic fragments are suitably labeled, such as by end-labeling with ³²PdATP using T4 Kinase. Labeled fragments are then incubated with nuclear extracts of TM cells, +/- treatment with DEX (500nM) for 10 days, and HeLa cell extract (Stratagene, CA) for 15 minutes at 0°C. The samples are then run on a low ionic strength, non-denaturing polyacrylamide gel. The gel is then dried and exposed to X-ray film overnight. The shift in mobility, apparent in Figure 9, demonstrates

the presence of DNA binding components from HTM cells, and that treatment with DEX changes the amount or strength of binding.

WHAT IS CLAIMED IS:

1. A method for diagnosing glaucoma in a patient which comprises the steps:

- (A) incubating under conditions permitting nucleic acid hybridization: a marker nucleic acid molecule, said first marker nucleic acid molecule comprising a nucleotide sequence of a polynucleotide that specifically hybridizes to a polynucleotide that is linked to a TIGR promoter, and a complementary nucleic acid molecule obtained from a cell or a bodily fluid of said patient, wherein nucleic acid hybridization between said marker nucleic acid molecule, and said complementary nucleic acid molecule obtained from said patient permits the detection of a polymorphism whose presence is predictive of a mutation affecting TIGR response in said patient;
- (B) permitting hybridization between said marker nucleic acid molecule and said complementary nucleic acid molecule obtained from said patient; and
- (C) detecting the presence of said polymorphism, wherein the detection of said polymorphism is diagnostic of glaucoma.
- 2. A method for diagnosing glaucoma in a patient according to claim 1, wherein said marker nucleic acid molecule is capable of specifically detecting *TIGRmt1*.
- 3. A method for diagnosing glaucoma in a patient according to claim 1, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRmt2.
- 4. A method for diagnosing glaucoma in a patient according to claim 1, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRmt3.
- 5. A method for diagnosing glaucoma in a patient according to claim 1, wherein said marker nucleic acid molecule is capable of specifically detecting *TIGRmt4*.
- 6. A method for diagnosing glaucoma in a patient according to claim 1, wherein said marker nucleic acid molecule is capable of specifically detecting *TIGRmt5*.
- 7. A method for diagnosing glaucoma in a patient according to claim 1, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRsv1.
- 8. A method for diagnosing glaucoma in a patient according to claim 1, further comprising a second marker nucleic acid molecule.
- 9. A method for diagnosing glaucoma in a patient according to claim 8, wherein said first marker nucleic acid molecule and said second marker nucleic acid molecule are selected from the group consisting of a nucleic acid molecule that comprises the sequence of SEQ ID NO: 6, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 7, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 8, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 9, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 10, a

nucleic acid molecule that comprises the sequence of SEQ ID NO: 11, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 12, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 14, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 16, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 17, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 18, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 19, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 20, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 21, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 23, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 23, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 24 and a nucleic acid molecule that comprises the sequence of SEQ ID NO: 25.

- 10. A method for diagnosing glaucoma in a patient according to claim 9, wherein said first marker nucleic acid molecule and said second marker nucleic acid molecule are selected from the group consisting of a nucleic acid molecule that comprises the sequence of SEQ ID NO: 6, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 7, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 8, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 9, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 12, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 18, and a nucleic acid molecule that comprises the sequence of SEQ ID NO: 25
- 11. A method for diagnosing glaucoma in a patient according to claim 10, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 12.
- 12. A method for diagnosing glaucoma in a patient according to claim 10, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 9 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 8.
- 13. A method for diagnosing glaucoma in a patient according to claim 10, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 7 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 6.
- 14. A method for diagnosing glaucoma in a patient according to claim 10, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID

NO: 18 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 25.

- 15. A method for diagnosing steroid sensitivity in a patient which comprises the steps:
- (A) incubating under conditions permitting nucleic acid hybridization: a marker nucleic acid molecule, said marker nucleic acid molecule comprising a nucleotide sequence of a polynucleotide that is linked to a TIGR promoter, and a complementary nucleic acid molecule obtained from a cell or a bodily fluid of said patient, wherein nucleic acid hybridization between said marker nucleic acid molecule, and said complementary nucleic acid molecule obtained from said patient permits the detection of a polymorphism whose presence is predictive of a mutation affecting TIGR response in said patient;
- (B) permitting hybridization between said TIGR-encoding marker nucleic acid molecule and said complementary nucleic acid molecule obtained from said patient; and
- (C) detecting the presence of said polymorphism, wherein the detection of said polymorphism is diagnostic of steroid sensitivity.
- 16. A method for diagnosing steroid sensitivity in a patient according to claim 15, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRmt1.
- 17. A method for diagnosing steroid sensitivity in a patient according to claim 15, wherein said marker nucleic acid molecule is capable of specifically detecting *TIGRmt2*.
- 18. A method for diagnosing steroid sensitivity in a patient according to claim 15, wherein said marker nucleic acid molecule is capable of specifically detecting *TIGRmt3*.
- 19. A method for diagnosing steroid sensitivity in a patient according to claim 15, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRmt4.
- 20. A method for diagnosing steroid sensitivity in a patient according to claim 15, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRmt5.
- 21. A method for diagnosing steroid sensitivity in a patient according to claim 15, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRsv1.
- 22. A method for diagnosing steroid sensitivity in a patient according to claim 15, further comprising a second marker nucleic acid molecule.
- 23. A method for diagnosing steroid sensitivity in a patient according to claim 22, wherein said first marker nucleic acid molecule and said second marker nucleic acid molecule are selected from the group consisting of a nucleic acid molecule that comprises the sequence of SEQ ID NO: 6, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 7, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 8, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 9, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 10, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 11, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 11, a nucleic acid molecule that comprises

the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 16, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 16, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 17, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 19, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 19, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 20, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 21, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 23, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 24 and a nucleic acid molecule that comprises the sequence of SEQ ID NO: 25.

- 24. A method for diagnosing steroid sensitivity in a patient according to claim 23, wherein said first marker nucleic acid molecule and said second marker nucleic acid molecule are selected from the group consisting of a nucleic acid molecule that comprises the sequence of SEQ ID NO: 6, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 7, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 8, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 9, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 12, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 18, and a nucleic acid molecule that comprises the sequence of SEQ ID NO: 25.
- 25. A method for diagnosing steroid sensitivity in a patient according to claim 24, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 12.
- 26. A method for diagnosing glaucoma in a patient according to claim 24, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 9 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 5.
- 27. A method for diagnosing steroid sensitivity in a patient according to claim 24, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 7 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 6.
- 28. A method for diagnosing steroid sensitivity in a patient according to claim 24, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 18 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 25.

29. The method of claims 10 or 24, wherein said complementary nucleic acid molecule obtained from a cell or a bodily fluid of said patient has been amplified using a nucleic acid amplification method.

- 30. The method of claim 1, wherein said marker nucleic acid molecule is selected from the group consisting of D1S2536 marker nucleic acid, D1S210 marker nucleic acid, D1S1552 marker nucleic acid, D1S2536 marker nucleic acid D1S2790 marker nucleic acid, SHGC-12820 marker nucleic acid, and D1S2558 marker nucleic acid.
- 31. The method of claim 30, wherein said marker nucleic acid molecule is D1S2536 marker nucleic acid.
- 32. The method of claim 15, wherein said marker nucleic acid molecule is selected from the group consisting of D1S2536 marker nucleic acid, D1S210 marker nucleic acid, D1S1552 marker nucleic acid, D1S2536 marker nucleic acid D1S2790 marker nucleic acid, SHGC-12820 marker nucleic acid, and D1S2558 marker nucleic acid.
- 33. The method of claim 32, wherein said marker nucleic acid molecule is D1S2536 marker nucleic acid.
- 34. A nucleic acid molecule that comprises the sequence of SEQ ID NO: 1.
- 35. A recombinant DNA molecule containing a polynucleotide that specifically hybridizes to SEQ ID NO: 1.
- 36. A substantially purified molecule that specifically binds to a nucleic acid molecule that comprises the sequence of SEQ ID NO:1.
- 37. A nucleic acid molecule that comprises the sequence of SEQ ID NO: 3.
- 38. A recombinant DNA molecule containing a polynucleotide that specifically hybridizes to SEQ ID NO: 3.
- 39. A substantially purified molecule that specifically binds to a nucleic acid molecule that comprises the sequence of SEQ ID NO: 3.
- 40. A nucleic acid molecule that comprises the sequence of SEQ ID NO: 4.
- 41. A recombinant DNA molecule containing a polynucleotide that specifically hybridizes to SEO ID NO: 4.
- 42. A substantially purified molecule that specifically binds to a nucleic acid molecule that comprises the sequence of SEQ ID NO: 4.
- 43. A nucleic acid molecule that comprises the sequence of SEQ ID NO: 5.
- 44. A recombinant DNA molecule containing a polynucleotide that specifically hybridizes to SEQ ID NO: 5.
- 45. A substantially purified molecule that specifically binds to a nucleic acid molecule that comprises the sequence of SEQ ID NO: 5.
- 46. A nucleic acid molecule that comprises the sequence of SEQ ID NO: 26.

47. A recombinant DNA molecule containing a polynucleotide that specifically hybridizes to SEQ ID NO: 26.

- 48. A substantially purified molecule that specifically binds to a nucleic acid molecule that comprises the sequence of SEQ ID NO: 26.
- A substantially purified molecule that specifically binds to a nucleic acid molecule 49. selected from the group consisting of a nucleic acid molecule that comprises a cis element characteristic of PRL-FP111, a nucleic acid molecule that comprises a glucocorticoid response cis element, a nucleic acid molecule that comprises a cis element characteristic of GR/PR, a nucleic acid molecule that comprises a shear stress response cis element, a nucleic acid molecule that comprises a glucocorticoid response cis element, a nucleic acid molecule that comprises a cis element characteristic of CBE, a nucleic acid molecule that comprises a cis element capable of binding NFE, a nucleic acid molecule that comprises a cis element capable of binding KTF.1-CS, a nucleic acid molecule that comprises a cis element characteristic of PRE, a nucleic acid molecule that comprises a cis element characteristic of ETF-EGFR, a nucleic acid molecule that comprises a cis element capable of binding SRE-cFos, a nucleic acid molecule that comprises a cis element characteristic of Alu, a nucleic acid molecule that comprises a cis element capable of binding VBP, a nucleic acid molecule that comprises a cis element characteristic of Malt-CS, a nucleic acid molecule that comprises a cis element capable of binding ERE, a nucleic acid molecule that comprises a cis element characteristic of NF-mutagen, a nucleic acid molecule that comprises a cis element capable of binding myc-PRF, a nucleic acid molecule that comprises a cis element capable of binding AP2, a nucleic acid molecule that comprises a cis element capable of binding HSTF, a nucleic acid molecule that comprises a cis element characteristic of SBF, a nucleic acid molecule that comprises a cis element capable of binding NF-1, a nucleic acid molecule that comprises a cis element capable of binding NF-MHCIIA/B, a nucleic acid molecule that comprises a cis element capable of binding PEA1, a nucleic acid molecule that comprises a cis element characteristic of ICS, a nucleic acid molecule that comprises a cis element capable of binding ISGF2, a nucleic acid molecule that comprises a cis element capable of binding zinc, a nucleic acid molecule that comprises a cis element characteristic of CAP/CRP-galO, a nucleic acid molecule that comprises a cis element capable of binding AP1, a nucleic acid molecule that comprises a cis element capable of binding SRY,, a nucleic acid molecule that comprises a cis element characteristic of GC2, a nucleic acid molecule that comprises a cis element capable of binding PEA3, a nucleic acid molecule that comprises a cis element characteristic of MIR, a nucleic acid molecule that comprises a cis element capable of binding NF-HNF-1, a nucleic acid molecule that comprises a thyroid receptor cis element, and a nucleic acid molecule that comprises a cis element capable of binding NFKB.

50. A method of treating glaucoma which comprises administering to a glaucomatous patient an effective amount of an agent capable of binding a cis element located within SEQ ID NO: 1.

- 51. The method of claim 50, wherein said agent inhibits the expression of a TIGR mRNA.
- 52. The method of claim 50, wherein said agent binds a DNA sequence within SEQ ID NO:
 1.

The method of claim 50, wherein said agent binds a nucleic acid molecule that comprises 53. element characteristic of PRL-FP111, a nucleic acid molecule that comprises a glucocorticoid response cis element, a nucleic acid molecule that comprises a cis element characteristic of GR/PR, a nucleic acid molecule that comprises a shear stress response cis element, a nucleic acid molecule that comprises a glucocorticoid response cis element, a nucleic acid molecule that comprises a cis element characteristic of CBE, a nucleic acid molecule that comprises a cis element capable of binding NFE, a nucleic acid molecule that comprises a cis element capable of binding KTF.1-CS, a nucleic acid molecule that comprises a cis element characteristic of PRE, a nucleic acid molecule that comprises a cis element characteristic of ETF-EGFR, a nucleic acid molecule that comprises a cis element capable of binding SRE-cFos, a nucleic acid molecule that comprises a cis element characteristic of Alu, a nucleic acid molecule that comprises a cis element capable of binding VBP, a nucleic acid molecule that comprises a cis element characteristic of Malt-CS, a nucleic acid molecule that comprises a cis element capable of binding ERE, a nucleic acid molecule that comprises a cis element characteristic of NF-mutagen, a nucleic acid molecule that comprises a cis element capable of binding myc-PRF, a nucleic acid molecule that comprises a cis element capable of binding AP2, a nucleic acid molecule that comprises a cis element capable of binding HSTF, a nucleic acid molecule that comprises a cis element characteristic of SBF, a nucleic acid molecule that comprises a cis element capable of binding NF-1, a nucleic acid molecule that comprises a cis element capable of binding NF-MHCIIA/B, a nucleic acid molecule that comprises a cis element capable of binding PEA1, a nucleic acid molecule that comprises a cis element characteristic of ICS, a nucleic acid molecule that comprises a cis element capable of binding ISGF2, a nucleic acid molecule that comprises a cis element capable of binding zinc, a nucleic acid molecule that comprises a cis element characteristic of CAP/CRP-galO, a nucleic acid molecule that comprises a cis element capable of binding AP1, a nucleic acid molecule that comprises a cis element capable of binding SRY,, a nucleic acid molecule that comprises a cis element characteristic of GC2, a nucleic acid molecule that comprises a cis element capable of binding PEA3, a nucleic acid molecule that comprises a cis element characteristic of MIR, a nucleic acid molecule that comprises a cis element capable of binding NF-HNF-1, a nucleic acid molecule that comprises a thyroid receptor cis element, and a nucleic acid molecule that comprises a cis element capable of binding NFkB.

- 54. A method for prognosing glaucoma in a patient which comprises the steps:
- (A) incubating under conditions permitting nucleic acid hybridization: a marker nucleic acid molecule, said first marker nucleic acid molecule comprising a nucleotide sequence of a polynucleotide that specifically hybridizes to a polynucleotide that is linked to a TIGR promoter, and a complementary nucleic acid molecule obtained from a cell or a bodily fluid of said patient, wherein nucleic acid hybridization between said marker nucleic acid molecule, and said complementary nucleic acid molecule obtained from said patient permits the detection of a polymorphism whose presence is predictive of a mutation affecting TIGR response in said patient;
- (B) permitting hybridization between said marker nucleic acid molecule and said complementary nucleic acid molecule obtained from said patient; and
- (C) detecting the presence of said polymorphism, wherein the detection of said polymorphism is prognostic of glaucoma.
- 55. A method for prognosing glaucoma in a patient according to claim 54, wherein said marker nucleic acid molecule is capable of specifically detecting *TIGRmt1*.
- 56. A method for prognosing glaucoma in a patient according to claim 54, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRmt2.
- 57. A method for prognosing glaucoma in a patient according to claim 54, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRmt3.
- 58. A method for prognosing glaucoma in a patient according to claim 54, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRmt4.
- 59. A method for prognosing glaucoma in a patient according to claim 54, wherein said marker nucleic acid molecule is capable of specifically detecting *TIGRmt5*.
- 60. A method for prognosing glaucoma in a patient according to claim 54, wherein said marker nucleic acid molecule is capable of specifically detecting TIGRsv1.
- 61. A method for prognosing glaucoma in a patient according to claim 54, further comprising a second marker nucleic acid molecule.
- 62. A method for prognosing glaucoma in a patient according to claim 61, wherein said first marker nucleic acid molecule and said second marker nucleic acid molecule are selected from the group consisting of a nucleic acid molecule that comprises the sequence of SEQ ID NO: 6, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 7, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 8, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 9, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 10, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 11, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 12, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 14, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecul

NO: 14, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 15, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 16, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 17, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 18, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 19, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 20, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 21, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 23, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 24 and a nucleic acid molecule that comprises the sequence of SEQ ID NO: 25.

- 63. A method for diagnosing glaucoma in a patient according to claim 62, wherein said first marker nucleic acid molecule and said second marker nucleic acid molecule are selected from the group consisting of a nucleic acid molecule that comprises the sequence of SEQ ID NO: 6, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 7, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 8, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 9, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 12, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13, a nucleic acid molecule that comprises the sequence of SEQ ID NO: 18, and a nucleic acid molecule that comprises the sequence of SEQ ID NO: 25
- 64. A method for diagnosing glaucoma in a patient according to claim 63, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 13 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 12.
- 65. A method for diagnosing glaucoma in a patient according to claim 63, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 9 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 8.
- 66. A method for diagnosing glaucoma in a patient according to claim 63, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 7 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 6.
- 67. A method for diagnosing glaucoma in a patient according to claim 63, wherein said first marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 18 and said second marker nucleic acid molecule is a nucleic acid molecule that comprises the sequence of SEQ ID NO: 25.
- 68. The method of claim 54, wherein said marker nucleic acid molecule is selected from the group consisting of D1S2536 marker nucleic acid, D1S210 marker nucleic acid, D1S1552

marker nucleic acid, D1S2536 marker nucleic acid D1S2790 marker nucleic acid, SHGC-12820 marker nucleic acid, and D1S2558 marker nucleic acid.

- 69. The method of claim 68, wherein said marker nucleic acid molecule is D1S2536 marker nucleic acid.
- 70. A nucleic acid comprising a nucleotide sequence selected from the group consisting of SEQ ID NO: 33 and its complement, a region of SEQ ID NO: 33 or its complement that specifically hybridizes to a nucleic acid possessing the characteristic C to T substitution of the mt11 sequence variant, and a region of SEQ ID NO: 33 or its complement that specifically hybridizes to a nucleic acid possessing the characteristic C to T substitution of the TIGRmt11 sequence variant but does not specifically hybridize to a nucleic acid that does not possess the TIGRmt11 sequence variant under high stringency conditions.
- 71. A nucleic acid that specifically hybridizes to the nucleic acid of claim 70.
- 72. A vector comprising the nucleic acid of claim 70.
- 73. A cell comprising the nucleic acid of claim 70.
- 74. A method for detecting the presence or absence of the characteristic TIGRmt11 sequence variation in a sample containing DNA, comprising contacting a labeled nucleic acid of claim 70 with the DNA of the sample under hybridization conditions and determining the presence of hybrid nucleic acid molecules comprising the labeled nucleic acid.
- 75. A method for determining the presence of increased susceptibility to a glaucoma, to a progressive ocular hypertensive disorder resulting in loss of visual field, or the presence of steroid sensitivity in a patient, comprising the method of claim 74, wherein the sample containing DNA is derived from the patient.
- 76. The method of claim 75, which is performed during or after the patient is treated with a steroid compound.
- 77. The method of claim 75, which is performed prior to an administration of a steroid compound.
- 78. A kit for determining the presence of increased susceptibility to a glaucoma, to a progressive ocular hypertensive disorder resulting in loss of visual field, or the presence of

steroid sensitivity in a patient, comprising a labeled nucleic acid of claim 70 and a means for detecting hybridization with the labeled nucleic acid.

- 79. A nucleic acid comprising a nucleotide sequence selected from the group consisting of one of SEQ ID NO: 1-3 or 34, and a fragment of SEQ ID NO: 1-3, or 34 that possesses a functional regulatory region.
- 80. A cell comprising an introduced nucleic acid of the sequence as claimed in claim 79.
- 81. A vector comprising a nucleic acid as claimed in claim 79.
- 82. A method for detecting the specific binding of a molecule to a nucleic acid comprising providing a nucleic acid of claim 79, contacting the nucleic acid with a sample containing the molecule to be tested, and identifying binding of the molecule to the nucleic acid.
- 83. A method as claimed in claim 82, wherein the identifying step comprises a gel shift assay.
- 84. A method as claimed in claim 82, wherein the nucleic acid is labeled.
- 85. A method for detecting the presence of the TIGRmt11 sequence variation in a sample containing DNA, comprising providing amplification reaction primers that direct amplification of a selected nucleic acid region containing the T to C substitution of the TIGRmt11 sequence variant, amplifying the nucleic acid defined by the amplification reaction primers, and determining the presence or absence of the T to C substitution in the amplified nucleic acid.
- 86. The method of claim 85, wherein the determining the presence or absence of the T to C substitution comprises sequencing the amplified nucleic acid.
- 87. The method of claim 86, wherein the determining the presence or absence of the T to C substitution comprises a hybridization assay.
- 88. A method for determining the presence of increased susceptibility to a glaucoma, to a progressive ocular hypertensive disorder resulting in loss of visual field, or the presence of steroid sensitivity in a patient comprising the method of claim 85, wherein the sample containing DNA is derived from the patient.
- 89. A kit for determining the presence of increased susceptibility to a glaucoma, to a progressive ocular hypertensive disorder resulting in loss of visual field, or the presence of steroid sensitivity in a patient, comprising amplification reaction primers that direct amplification of a selected nucleic acid region containing the T to C substitution of the TIGRmt11 sequence variant and an enzyme for amplifying the region containing the T to C substitution.
- 90. A method for detecting a polymorphism in the 5' flanking region of a TIGR gene, comprising selecting amplification reaction primers from the group consisting of nucleic acids comprising nucleotide sequences SEQ ID NO: 6-25 or 35, or complements thereof, nucleotide sequences from a fragment of SEQ ID NO: 6-25 or 35, or their complements, and nucleotide sequences from an about 18 to an about 60 nucleotide fragment of the 5' flanking sequences in

SEQ ID NO: 1-3, or 34, or complements thereof, amplifying a selected nucleic acid region of the 5' flanking region defined by the amplification reaction primers in a sample of DNA, and comparing at least part of the sequence of the amplified nucleic acid with the sequence set forth in SEQ ID NO: 1-3.

- 91. A substantially purified nucleic acid molecule comprising SEQ ID NO: 37 or SEQ ID NO: 38.
- 92. The nucleic acid molecule of claim 91 that is capable of conferring tissue specific promoter activity.
- 93. A recombinant nucleic acid molecule comprising the nucleic acid molecule of claim 91.
- 94. A vector comprising a nucleic acid molecule of claim 91.
- 95. A cell comprising an introduced nucleic acid molecule, the introduced nucleic acid molecule comprising the nucleic acid molecule of claim 91.
- 96. A substantially purified nucleic acid molecule comprising a sequence having about 95% identity to SEQ ID NO: 37 or SEQ ID NO: 38.
- 97. The nucleic acid molecule of claim 96 that is capable of conferring a tissue specific promoter activity.
- 98. A recombinant nucleic acid molecule comprising the nucleic acid molecule of claim 96.
- 99. A vector comprising a nucleic acid molecule of claim 96.
- 100. A cell comprising an introduced nucleic acid molecule, the introduced nucleic acid molecule comprising the nucleic acid molecule of claim 96.
- 101. A substantially purified nucleic acid molecule comprising SEQ ID NO: 37 or SEQ ID NO: 38, or a variant thereof, capable of conferring tissue specific expression.
- 102. A recombinant nucleic acid molecule comprising the nucleic acid molecule of claim 101.
- 103. A vector comprising a nucleic acid molecule of claim 101.
- 104. A cell comprising an introduced nucleic acid molecule, the introduced nucleic acid molecule comprising the nucleic acid molecule of claim 101.
- 105. A method for identifying a protein or first compound that binds to a sequence of a TIGR gene, comprising incubating a nucleic acid molecule comprising SEQ ID NO: 37 or 38, or variant or either, or a region of SEQ ID NO: 3 or 34, with a composition containing the protein or first compound, and detecting the presence of binding to the nucleic acid molecule.

106. A method for identifying a second compound that modulates the binding of a protein or first compound to a TIGR gene sequence, comprising the steps of claim 105, further comprising adding the second compound to the nucleic acid molecule and comparing the binding detected with a control.

- 107. A method for identifying a protein or compound that modulates expression of a TIGR gene, comprising incubating a nucleic acid molecule comprising SEQ ID NO: 37 or 38, or variant of either, or a region of SEQ ID NO: 3 or 34, with a composition containing the protein or compound, and detecting the presence of binding to the nucleic acid molecule compared to a control.
- 108. The method of claim 107 wherein the nucleic acid molecule is contained within a cell.
- 109. A method for identifying a cellular component that modulates expression of a TIGR gene, comprising incubating a nucleic acid molecule comprising SEQ ID NO: 37 or 38, or variant of either, or a region of SEQ ID NO: 3 or 34, with an extract of a cell, and detecting the presence of specific binding to the nucleic acid molecule.
- 110. A method for identifying a compound that modulates steroid induction of a TIGR gene, comprising incubating a cell containing an introduced nucleic acid molecule comprising a TIGR 5'regulatory region with the compound, and detecting the expression of a gene linked to the TIGR 5' regulatory region.
- 111. The method of claim 110, wherein the 5" regulatory sequence comprises SEQ ID NO: 37 or 38, or variant of either, or a region of SEQ ID NO: 3 or 34.

1 ATC TITGTTCAGT TTACCTCAGG GCTATTATGA 33	
34 AATGAAATGA GATAACCAAT GTGAAAGTCC TATAAACTGT ATAGCCTCCA TTCGGATGTA 93	
94 TGTCTTTGGC AGGATGATAA AGAATCAGGA AGAAGGAGTA TCCACGTTAG CCAAGTGTCC 153	
154 AGGCTGTGTC TGCTCTTATT TTAGTGACAG ATGTTGCTCC TGACAGAAGC TATTCTTCAG 213	
214 GAAACATCAC ATCCAATATG GTAAATCCAT CAAACAGGAG CTAAGAAACA GGAATGAGAT 273	
274 GGGCACTTGC CCAAGGAAAA ATGCCAGGAG AGCAAATAAT GATGAAAAAT AAACTTTTCC 333	
334 CTTTGTTTTT AATTTCAGGA AAAAATGATG AGGACCAAAA TCAATGAATA AGGAAAACAG 393 (Prl.FPIII) CCTG AAAATGAATA AGAAA	
394 CTCAGAAAAA AGATGTTTCC AAATTGGTAA TTAAGTATTT GTTCCTTGGG AAGAGACCTC 453 (PR/GR-MMTV) T GTTCTTTTGG AA (SSRE) GAGACC	
454 CATGTGAGCT TGATGGGAAA ATGGGAAAAA CGTCAAAAGC ATGATCTGAT CAGATCCCAA 513	
514 AGTGGATTAT TATTTTAAAA ACCAGATGGC ATCACTCTGG GGAGGCAAGT TCAGGAAGGT 573	
574 CATGTTAGCA AAGGACATAA CAATAACAGC AAAATCAAAA TTCCGCAAAT GCAGGAGGAA 633 CCTTTTAG-A AAGGACAAAA CAGAATG (nGRE-PRL)	
634 AATGGGGACT GGGAAAGCTT TCATAACAGT GATTAGGCAG TTGACCATGT TCGCAACACC 693	
694 TCCCCGTCTA TACCAGGGAA CACAAAAATT GACTGGGCTA AGCCTGGACT TTCAAGGGAA 753 GCCTGGACT GTC (CBE-P53)	
754 ATATGAAAAA CTGAGAGCAA AACAAAAGAC ATGGTTAAAA GGCAACCAGA ACATTGTGAG 813 ATTTTTCTGA TTGGTTAAAA GT (NFEi)	
814 CCTTCAAAGC AGCAGTGCCC CTCAGCAGGG ACCCTGAGGC ATTTGCCTTT AGGAAGGCCA 873 G ACCCTGAGGC T (KTF.1-CS)	
874 GTTTTCTTAA GGAATCTTAA GAAACTCTTG AAAGATCATG AATTTTAACC ATTTTAAGTA 933	i
934 TAAAACAAAT ATGCGATGCA TAATCAGTTT AGACATGGGT CCCAATTTTA TAAAGTCAGG 993 (PRE-lysozyme) AGGCCGT	1
994 CATACAAGGA TAACGTGTCC CAGCTCCGGA TAGGTCAGAA ATCATTAGAA ATCACTGTGT 105 GATCCAAGGA GCAGAAGTTC CAGCTATGGT CAG (GRE-hMT) GG TACACTGTGT	3
1054 CCCCATCCTA ACTITITCAG AATGATCTGT CATAGCCCTC ACACACAGGC CCGATGTGTC 11	L13
1114 TGACCTACAA CCACATCTAC AACCCAAGTG CCTCAACCAT TGTTAACGTG TCATCTCAGT 11	173

FIG.1A

- 174 AGGTCCCATT ACAAATGCCA CCTCCCCTGT GCAGCCCATC CCGCTCCACA GGAAGTCTCC 1233

 1234 CCACTCTAGA CTTCTGCATC ACGATGTTAC AGCCAGAAGC TCCGTGAGGG TGAGGGTCTG 1293

 1294 TGTCTTACAC CTACCTGTAT GCTCTACACC TGAGCTCACT GCAACCTCTG CCTCCCAGGT 1353

 1354 TCAAGCAATT CTCCTGTCTC AGCCTCCCGC GTAGCTGGGA CTACAGGCGC ACGCCCGGCT 1413

 1414 AATTTTGTA TTGTTAGTAG AGATGGGGTT TCACCATATT AGCCCGGCTG GTCTTGAACT 1473

 Alu Repeat Region CCACCTCAGC CTCCTAAAGT GCTGGGATTA CAGGCATGAG 1533

 1534 TCACCGCGCC CGGCCAAGGG TCAGTGTTTA ATAAGGAATA ACTTGAATGG TTTACTAAAC 1593

 1594 CAACAGGGAA ACAGACAAAA GCTGTGATAA TTTCAGGGAT TCTTGGGATG GGGAATGGTG 1653

 1654 CCATGAGCTG CCTGCCTAGT CCCAGACCAC TGGTCCTCAT CACTTTCTC CCTCATCCTC 1713
 - 1834 GGAGGAGGC ATACCCCAGA GACTCCTTGA AGCCCCCGGC AGAGGTTTCC TCTCCAGCTG 1893 GGAKGAGG (Mait-cs)

1774 ACTGAAATAA GAGTATACAT AAACTAGTTC CATTTGGGGC CATCTGTGTG TGTGTATAGG 1833

GTTTACAT AAAC (VBP-vitel)

- 1894 GGGGAGCCCT GCAAGCACCC GGGGTCCTGG GTGTCCTGAG CAACCTGCCA GCCCGTGCCA 1953
- 1954 CTGGTTGTTT TGTTATCACT CTCTAGGGAC CTGTTGCTTT CTATTTCTGT GTGACTCGTT 2013
- 2014 CATTCATCCA GGCATTCATT GACAATTTAT TGAGTACTTA TATCTGCCAG ACACCAGAGA 2073
- 2074 CAAAATGGTG AGCAAAGCAG TCACTGCCCT ACCTTCGTGG AGGTGACAGT TTCTCATGGA 2133
- 2134 AGACGTGCAG AAGAAAATTA ATAGCCAGCC AACTTAAACC CAGTGCTGAA AGAAAGGAAA 2193 GCGTGAC CGGAGCTGAA AGAAAGGAAC
- 2194 TAAACACCAT CTTGAAGAAT TGTGCGCAGC ATCCCTTAAC AAGGCCACCT CCCTAGCGCC 2253 AC (ERE-c.vitel)
- 2254 CCCTGCTGCC TCCATCGTGC CCGGAGGCCC CCAAGCCCGA GTCTTCCAAG CCTCCTCCTC 2313
- 2314 CATCAGTCAC AGCGCTGCAG CTGGCCTGCC TCGCTTCCcG TGAATCGTCC TGGTGCATCT 2373
 AGCAG CTGGC (NF-mutagen)
- 2374 GAGCTGGAGA CTCCTTGGCT CCAGGCTCCA GAAAGGAAAT GGAGAGGGAA ACTAGTCTAA 2433 A GAAAGGGAAA GGA (PRF-myc)
- 2434 CGGAGAATCT GGAGGGGACA GTGTTTCCTC AGAGGGAAAG GGGCCTCCAC GTCCAGGAGA 2493
 ACCCGGTACA CTGTGTCCTC CCGCT (GRE-hMT.IIa)
 CC CTTTGGGCCA ATGTGTCCTG AGGGGA (GRE-hGH)

FIG.1B

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2494 ATTCCAGGAG GTGGGGACTG CAGGGAGTGG GGACGCTGGG GCTGAGCGGG TGCTGAAAGG 2553 CTGG GGAGCCTGGG GA (AP.2-SV40)

- 2554 CAGGAAGGTG AAAAGGGCAA GGCTGAAGCT GCCCAGATGT TCAGTGTTGT TCACGGGGCT 2613
- 2614 GGGAGTTTTC CGTTGCTTCC TGTGAGCCTT TTTATCTTTT CTCTGCTTGG AGGAGAAGAA 2673 CT CGTTGCTTCG AG (HSTF-hsp70)
- 2674 GTCTATTTCA TGAAGGGATG CAGTTTCATA AAGTCAGCTG TTAAAATTCC AGGGTGTGCA 2733
- 2734 TGGGTTTTCC TTCACGAAGG CCTTTATTTA ATGGGAATAT AGGAAGCGAG CTCATTTCCT 2793 TGGGTTTTTG (SBF.yeast)
- 2794 AGGCCGTTAA TTCACGGAAG AAGTGACTGG AGTCTTTTCT TTCATGTCTT CTGGGCAACT 2853
- 2854 ACTCAGCCCT GTGGTGGACT TGGCTTATGC AAGACGGTCG AAAACCTTGG AATCAGGAGA 2913
- 2914 CTCGGTTTTC TTTCTGGTTC TGCCATTGGT TGGCTGTGCG ACCGTGGGCA AGTGTCTCTC 2973
 C TTTCTGGTTT TGCAG (NF.1-bithorax)
 (NF-MHCII/)CCATTGGT T
- 2974 CTTCCCTGGG CCATAGTCTT CTCTGCTATA AAGACCCTTG CAGCTCTCGT GTTCTGTGAA 3033
- 3034 CACTTCCCTG TGATTCTCTG TGAGGGGGGA TGTTGAGAGG GGAAGGAGGC AGAGCTGGAG 3093
- 3094 CAGCTGAGCC ACAGGGAGG TGGAGGGGGA CAGGAAGGCA GGCAGAAGCT GGGTGCTCCA 3153
- 3154 TCAGTCCTCA CTGATCACGT CAGACTCCAG GACCGAGAGC CACAATGCTT CAGGAAAGCT 2943
- 2944 CAATGAACCC AACAGCCACA TTTTCCTTCC CTAAGCATAG ACAATGGCAT TTGCCAATAA 3273
- 3274 CCAAAAAGAA TGCAGAGACT AACTGGTGGT AGCTTTTGCC TGGCATTCAA AAACTGGGCC 3333 GAAGTGACT AACTG (PEA.1-Polyoma)
- 3334 AGAGCAAGTG GAAAATGCCA GAGATTGTTA AACTTTTCAC CCTGACCAGC ACCCCACGCA 3393
- 3394 GCTCAGCAGT GACTGCTGAC AGCACGGAGT GACCTGCAGC GCAGGGGAGG AGAAGAAAAA 3453 C AGGTCAGAGT GACCTG (ERE.2-Vitel.)
- 3454 GAGAGGGATA GTGTATGAGC AAGAAAGACA GATTCATTCA AGGGCAGTGG GAATTGACCA 3513
- 3514 CAGGGATTAT AGTCCACGTG ATCCTGGGTT CTAGGAGGCA GGGCTATATT GTGGGGGGAA 3573 (GRE-FLV) CGGGATAC CGAGAGAACA GGGCTATAGG
- 3574 AAAATCAGTT CAAGGGAAGT CGGGAGACCT GATTTCTAAT ACTATATTTT TCCTTTACAA 3633 GAGACC (SSRE)
- 3634 GCTGAGTAAT TCTGAGCAAG TCACAAGGTA GTAACTGAGG CTGTAAGATT ACTTAGTTTC 3693 (ICS-MTII/ HLA-DR/)AGTTTC
- 3694 TCCTTATTAG GAACTCTTTT TCTCTGTGGA GTTAGCAGCA CAAGGGCAAT CCCGTTTCTT 3753
 TCCTCT
- 3754 TTAACAGGAA GAAAACATTC CTAAGAGTAA AGCCAAACAG ATTCAAGCCT AGGTCTTGCT 3813
- 3814 GACTATATGA TTGGTTTTTT GAAAAATCAT TTCAGCGATG TTTACTATCT GATTCAGAAA 3873

- 3874 ATGAGACTAG TACCCTTTGG TCAGCTGTAA ACAAACACCC ATTTGTAAAT GTCTCAAGTT 3933 GG TCA (1/2 ERE)
- 3934 CAGGCTTAAC TGCAGAACCA ATCAAATAAG AATAGAATCT TTAGAGCAAA CTGTGTTTCT 3993
- 3994 CCACTCTGGA GGTGAGTCTG CCAGGGCAGT TTGGAAATAT TTACTTCACA AGTATTGACA 4053
- 4054 CTGTTGTTGG TATTAACAAC ATAAAGTTGC TCAAAGGCAA TCATTATTTC AAGTGGCTTA 4113
- 4114 AAGTTACTTC TGACAGTTTT GGTATATTTA TTGGCTATTG CCATTTGCTT TTTGTTTTT 4173
 (NF.1-HCMV)TTGGCTATTG GCCA CTTT
- 4174 CTCTTTGGGT TTATTAATGT AAAGCAGGGA TTATTAACCT ACAGTCCAGA AAGCCTGTGA 4233 CTCTTT (ISGF2)
- 4234 ATTTGAATGA GGAAAAAATT ACATTTTTGT TTTTACCACC TTCTAACTAA ATTTAACATT 4293
 (Zn binding)------
- 4294 TTATTCCATT GCGAATAGAG CCATAAACTC AAAGTGGTAA TAACAGTACC TGTGATTTTG 4353
- 4354 TCATTACCAA TAGAAATCAC AGACATTTTA TACTATATTA CAGTTGTTGC AGATACGTTG 4413 (CAP-gal0) ATTTA TTCCATGTCA CACTTTTCGC A
- 4414 TAAGTGAAAT ATTTATACTC AAAACTACTT TGAAATTAGA CCTCCTGCTG GATCTTGTTT 4473
 TTACTC A (AP-1)
- 4474 TTAACATATT AATAAAACAT GTTTAAAATT TTGATATTIT GATAATCATA TTTCATTATC 4533
 GAT GTTTAAAAAT (PRL-FPII)
- 4534 ATTTGTTTCC TTTGTAATCT ATATTTTATA TATTTGAAAA CATCTTTCTG AGAAGAGTTC 4593 (GRE-MuRFV) TGTTTTCTG AGAACATCAG
- 4594 CCCAGATTTC ACCAATGAGG TTCTTGGCAT GCACACACAC AGAGTAAGAA CTGATTTAGA 4653
 CCAGATCTC ACCATCATTAT (nGRE) CACACACAC A (CACA)
 CTCTGG GGACAC AGAGTAGGG (AP.1-TGFb)
- 4654 GGCTAACATT GACATTGGTG CCTGAGATGC AAGACTGAAA TTAGAAAGTT CTCCCAAAGA 4713
 (GC2) GATGCT GATGGATAAT TTAGAAGCTT CTCCCACA
- 4714 TACACAGTTG TTTTAAAGCT AGGGGTGAGG GGGGAAATCT GCCGC<u>TTCTA TAGGAATGCT</u> 4773 (PEA.3)AGGAA GGT_
- 4774 CTCCCTGGAG CCTGGTAGGG TGCTGTCCTT GTGTTCTGGC TGGCTGTTAT TTTTCTCTGT 4833 CTC (SSRE) MIR Repeat Region
- 4834 CCCTGCTACG TCTTAAAGGA CTTGTTTGGA TCTCCAGTTC CTAGCATAGT GCCTGGCACA 4893

 GGA CTTGTTTGTT CT (GRE-rTAT-II) TGGGCACA

 GCAAAAAGGA TCTATTTGGA A (GRE-MMTV)
- 4894 <u>GTGCAGGTTC TCAATGAGTT TGCAGAGTGA ATGGAAATAT AAACTAGAAA TATATCCTTG</u> 4953 GTGCCAA (NF-1 (HNF-1)C TGTGAAATAT TAACTAAA
- 4954 TTGAAATCAG CACACCAGTA GTCCTGGTGT AAGTGTGTGT ACGTGTGTGT GTGTGTGTG 5013

FIG.1D

- 5014 GTGTGTGTT AAAACCAGGT GGAGATATAG GAACTATTAT TGGGGTATGG GTG<u>CATAAAT</u> 5073 cat/reverse cat box
- 5074 TGGGATGTTC TTTTTAAAAA GAAACTCCAA ACAGACTTCT GGAAGGTTAT TTTCTAAGAA 5133 (1/2GRE)TGTTC T (HSTF) GAAACTTCT GGAATATTCC CGAACTTTC CTTTTAGAAA GGA---CAAA ACAGAATG(nGRE-Pr1)
- 5134 TCTTGCTGGC AGCGTGAAGG CAACCCCCCT GTGCACAGCC CCACCCAGCC TCACGTGGCC 5193
 (1/2 TRE)AGG CAA T-CC CCAGGCTCCC -CAG(AP.2-SV40)
 GGAGAGCC CC (NF-KB)
- 5194 ACCTETGTCT TCCCCCATGA AGGGCTGGCT CCCCAG<u>TATA TATAAA</u>CCTC TCTGGAGCTC 5253 tata box GGTC TC (SSRE)
- 5254 GGGCATGAGC CAGCAAGGC*C* ACCCATCCAG GCACCTCTCA GCACAGC 5300 Start Sites

FIG.1E

ATC TITGTTCAGT TTACCTCAGG GCTATTATGA 33 1 34 AATGAAATGA GATAACCAAT GTGAAAGTCC TATAAACTGT ATAGCCTCCA TTCGGATGTA 93 94 TGTCTTTGGC AGGATGATAA AGAATCAGGA AGAAGGAGTA TCCACGTTAG CCAAGTGTCC 153 154 AGGCTGTGTC TGCTCTTATT TTAGTGACAG ATGTTGCTCC TGACAGAAGC TATTCTTCAG 213 214 GAAACATCAC ATCCAATATG GTAAATCCAT CAAACAGGAG CTAAGAAACA GGAATGAGAT 273 274 GGGCACTTGC CCAAGGAAAA ATGCCAGGAG AGCAAATAAT GATGAAAAAT AAACTTTTCC 333 334 CTTTGTTTTT AATTTCAGGA AAAAATGATG AGGACCAAAA TCAATGAATA AGGAAAACAG 393 394 CTCAGAAAAA AGATGTTTCC AAATTGGTAA TTAAGTATTT GTTCCTTGGG AAGAGACCTC 453 454 CATGTGAGCT TGATGGGAAA ATGGGAAAAA CGTCAAAAGC ATGATCTGAT CAGATCCCAA 513 514 AGTGGATTAT TATTTTAAAA ACCAGATGGC ATCACTCTGG GGAGGCAAGT TCAGGAAGGT 573 574 CATGTTAGCA AAGGACATAA CAATAACAGC AAAATCAAAA TTCCGCAAAT GCAGGAGGAA 633 634 AATGGGGACT GGGAAAGCTT TCATAACAGT GATTAGGCAG TTGACCATGT TCGCAACACC 693 694 TCCCCGTCTA TACCAGGGAA CACAAAAATT GACTGGGCTA AGCCTGGACT TTCAAGGGAA 753 754 ATATGAAAAA CTGAGAGCAA AACAAAAGAC ATGGTTAAAA GGCAACCAGA ACATTGTGAG 813 814 CCTTCAAAGC AGCAGTGCCC CTCAGCAGGG ACCCTGAGGC ATTTGCCTTT AGGAAGGCCA 873 874 GTTTTCTTAA GGAATCTTAA GAAACTCTTG AAAGATCATG AATTTTAACC ATTTTAAGTA 933 934 TAAAACAAAT ATGCGATGCA TAATCAGTTT AGACATGGGT CCCAATTTTA TAAAGTCAGG 993 994 CATACAAGGA TAACGTGTCC CAGCTCCGGA TAGGTCAGAA ATCATTAGAA ATCACTGTGT 1053 1054 CCCCATCCTA ACTITITCAG AATGATCTGT CATAGCCCTC ACACACAGGC CCGATGTGTC 1113 1114 TGACCTACAA CCACATCTAC AACCCAAGTG CCTCAACCAT TGTTAACGTG TCATCTCAGT 1173 1174 AGGTCCCATT ACAAATGCCA CCTCCCCTGT GCAGCCCATC CCGCTCCACA GGAAGTCTCC 1233 1234 CCACTCTAGA CTTCTGCATC ACGATGTTAC AGCCAGAAGC TCCGTGAGGG TGAGGGTCTG 1293 1294 TGTCTTACAC CTACCTGTAT GCTCTACACC TGAGCTCACT GCAACCTCTG CCTCCCAGGT 1353 1354 TCAAGCAATT CTCCTGTCTC AGCCTCCCGC GTAGCTGGGA CTACAGGCGC ACGCCCGGCT 1413 1414 AATTTTTGTA TTGTTAGTAG AGATGGGGTT TCACCATATT AGCCCGGCTG GTCTTGAACT 1473

FIG.2A

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1474 CCTGACCTCA GGTGATCCAC CCACCTCAGC CTCCTAAAGT GCTGGGATTA CAGGCATGAG 1533 1534 TCACCGCGCC CGGCCAAGGG TCAGTGTTTA ATAAGGAATA ACTTGAATGG TTTACTAAAC 1593 1594 CAACAGGGAA ACAGACAAAA GCTGTGATAA TTTCAGGGAT TCTTGGGATG GGGAATGGTG 1653 1654 CCATGAGCTG CCTGCCTAGT CCCAGACCAC TGGTCCTCAT CACTTTCTTC CCTCATCCTC 1713 1714 ATTITCAGGC TAAGTTACCA TITTATTCAC CATGCTTTTG TGGTAAGCCT CCACATCGTT 1773 1774 ACTGAAATAA GAGTATACAT AAACTAGTTC CATTTGGGGC CATCTGTGTG TGTGTATAGG 1833 1834 GGAGGAGGGC ATACCCCAGA GACTCCTTGA AGCCCCCGGC AGAGGTTTCC TCTCCAGCTG 1893 1894 GGGGAGCCCT GCAAGCACCC GGGGTCCTGG GTGTCCTGAG CAACCTGCCA GCCCGTGCCA 1953 1954 CTGGTTGTTT TGTTATCACT CTCTAGGGAC CTGTTGCTTT CTATTTCTGT GTGACTCGTT 2013 2014 CATTCATCCA GGCATTCATT GACAATTTAT TGAGTACTTA TATCTGCCAG ACACCAGAGA 2073 2074 CAAAATGGTG AGCAAAGCAG TCACTGCCCT ACCTTCGTGG AGGTGACAGT TTCTCATGGA 2133 2134 AGACGTGCAG AAGAAAATTA ATAGCCAGCC AACTTAAACC CAGTGCTGAA AGAAAGGAAA 2193 2194 TAAACACCAT CTTGAAGAAT TGTGCGCAGC ATCCCTTAAC AAGGCCACCT CCCTAGCGCC 2253 2254 CCCTGCTGCC TCCATCGTGC CCGGAGGCCC CCAAGCCCGA GTCTTCCAAG CCTCCTCCTC 2313 2314 CATCAGTCAC AGCGCTGCAG CTGGCCTGCC TCGCTTCCCG TGAATCGTCC TGGTGCATCT 2373 2374 GAGCTGGAGA CTCCTTGGCT CCAGGCTCCA GAAAGGAAAT GGAGAGGGAA ACTAGTCTAA 2433 2434 CGGAGAATCT GGAGGGGACA GTGTTTCCTC AGAGGGAAAG GGGCCTCCAC GTCCAGGAGA 2493 2494 ATTCCAGGAG GTGGGGACTG CAGGGAGTGG GGACGCTGGG GCTGAGCGGG TGCTGAAAGG 2553 2554 CAGGAAGGTG AAAAGGGCAA GGCTGAAGCT GCCCAGATGT TCAGTGTTGT TCACGGGGCT 2613 2614 GGGAGTTTTC CGTTGCTTCC TGTGAGCCTT TTTATCTTTT CTCTGCTTGG AGGAGAAGAA 2673 2674 GTCTATTTCA TGAAGGGATG CAGTTTCATA AAGTCAGCTG TTAAAATTCC AGGGTGTGCA 2733

3034 CACTTCCCTG TGATTCTCTG TGAGGGGGGA TGTTGAGAGG GGAAGGAGGC AGAGCTGGAG 3093 FIG.2B

2734 TGGGTTTTCC TTCACGAAGG CCTTTATTTA ATGGGAATAT AGGAAGCGAG CTCATTTCCT 2793

2794 AGGCCGTTAA TTCACGGAAG AAGTGACTGG AGTCTTTCT TTCATGTCTT CTGGGCAACT 2853

2854 ACTCAGCCCT GTGGTGGACT TGGCTTATGC AAGACGGTCG AAAACCTTGG AATCAGGAGA 2913

2914 CTCGGTTTTC TTTCTGGTTC TGCCATTGGT TGGCTGTGCG ACCGTGGGCA AGTGTCTCTC 2973

2974 CTTCCCTGGG CCATAGTCTT CTCTGCTATA AAGACCCTTG CAGCTCTCGT GTTCTGTGAA 3033

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3094 CAGCTGAGCC ACAGGGGAGG TGGAGGGGGA CAGGAAGGCA GGCAGAAGCT GGGTGCTCCA 3153 3154 TCAGTCCTCA CTGATCACGT CAGACTCCAG GACCGAGAGC CACAATGCTT CAGGAAAGCT 2943 2944 CAATGAACCC AACAGCCCACA TTTTCCTTCC CTAAGCATAG ACAATGGCAT TTGCCAATAA 3273 3274 CCAAAAAGAA TGCAGAGACT AACTGGTGGT AGCTTTTGCC TGGCATTCAA AAACTGGGCC 3333 3334 AGAGCAAGTG GAAAATGCCA GAGATTGTTA AACTTTTCAC CCTGACCAGC ACCCCACGCA 3393 3394 GCTCAGCAGT GACTGCTGAC AGCACGGAGT GACCTGCAGC GCAGGGGAGG AGAAGAAAAA 3453 3454 GAGAGGGATA GTGTATGAGC AAGAAAGACA GATTCATTCA AGGGCAGTGG GAATTGACCA 3513 3514 CAGGGATTAT AGTCCACGTG ATCCTGGGTT CTAGGAGGCA GGGCTATATT GTGGGGGGAA 3573 3574 AAAATCAGTT CAAGGGAAGT CGGGAGACCT GATTTCTAAT ACTATATTTT TCCTTTACAA 3633 3634 GCTGAGTAAT TCTGAGCAAG TCACAAGGTA GTAACTGAGG CTGTAAGATT ACTTAGTTTC 3693 3694 TCCTTATTAG GAACTCTTTT TCTCTGTGGA GTTAGCAGCA CAAGGGCAAT CCCGTTTCTT 3753 3754 TTAACAGGAA GAAAACATTC CTAAGAGTAA AGCCAAACAG ATTCAAGCCT AGGTCTTGCT 3813 3814 GACTATATGA TTGGTTTTTT GAAAAATCAT TTCAGCGATG TTTACTATCT GATTCAGAAA 3873 3874 ATGAGACTAG TACCCTTTGG TCAGCTGTAA ACAAACACCC ATTTGTAAAT GTCTCAAGTT 3933 3934 CAGGCTTAAC TGCAGAACCA ATCAAATAAG AATAGAATCT TTAGAGCAAA CTGTGTTTCT 3993 3994 CCACTCTGGA GGTGAGTCTG CCAGGGCAGT TTGGAAATAT TTACTTCACA AGTATTGACA 4053 4054 CTGTTGTTGG TATTAACAAC ATAAAGTTGC TCAAAGGCAA TCATTATTTC AAGTGGCTTA 4113 4114 AAGTTACTIC TGACAGTITT GGTATATITA TIGGCTATTG CCATTIGCTT TITGTTTTT 4173 4174 CTCTTTGGGT TTATTAATGT AAAGCAGGGA TTATTAACCT ACAGTCCAGA AAGCCTGTGA 4233 4234 ATTTGAATGA GGAAAAAATT ACGTTTTTAT TTTTACCACC TTCTAACTAA ATTTAACATT 4293 4294 TTATTCCATT GCGAATAGAG CCATAAACTC AAAGTGGTAA TAAGAGTACC TGTGATTTTG 4353 4354 TCATTACCAA TAGAAATCAC AGACATTTTA TACTATATTA CAGTTGTTGC AGGTACGTTG 4413 4414 TAAGTGAAAT ATTTATACTC AAAACTACTT TGAAATTAGA CCTCCTGCTG GATCTTGTTT 4473 4474 TTAACATATT AATAAAACAT GTTTAAAATT TTGATATTTT GATAATCATA TTTCATTATC 4533 4534 ATTTGTTTCC TTTGTAATCT ATATTTATA TATTTGAAAA CATCTTTCTG AGAAGAGTTC 4593 4594 CCCAGATTTC ACCAATGAGG TTCTTGGCAT GCACACACA AGAGTAAGAA CTGATTTAGA 4653 4654 GGCTAACATT GACATTGGTG CCTGAGATGC AAGACTGAAA TTAGAAAGTT CTCCCAAAGA 4713

FIG.2C

4714 TACACAGTTG TTTTAAAGCT AGGGGTGAGG GGGGAAATCT GCCGCTTCTA TAGGAATGCT 4773
4774 CTCCCTGGAG CCTGGTAGGG TGCTGTCCTT GTGTTCTGGC TGGCTGTTAT TTTTCTCTGT 4833
4834 CCCTGCTACG TCTTAAAGGA CTTGTTTGGA TCTCCAGTTC CTAGCATAGT GCCTGGCACA 4893
4894 GTGCAGGTTC TCAATGAGTT TGCAGAGTGA ATGGAAATAT AAACTAGAAA TATATCTTTG 4953
4954 TTGAAATCAG CACACCAGTA GTCCTGGTGT AAGTGTGTGT ACGTGTGTGTGTGT GTGTGTGTGT5017
5018 GTGTGTGTT AAAACCAGGT GGAGATATAG GAACTATTAT TGGGGTATGG GTGCATAAAT 5077
5078 TGGGATGTTC TTTTTAAAAA GAAACTCCAA ACAGACTTCT GGAAGGTTAT TTTCTAAGAA 5137
5138 TCTTGCTGGC AGCGTGAAGG CAACCCCCCT GTGCACAGCC CCACCCAGCC TCACGTGGCC 5197
5198 ACCTCTGTCT TCCCCCATGA AGGGCTGGCT CCCCAGTATA TATAAACCTC TCTGGAGCTC 5257
5258 GGGCATGAGC CAGCAAGGCC ACCCATCCAG GCACCTCTCA GCACAGC 5304

FIG.2D

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51 AATGTGAAAG TCCTATAAAC TGTATAGCCT CCATTCGGAT GTATGTCTTT 101 GGCAGGATGA TAAAGAATCA GGAAGAAGGA GTATCCACGT TAGCCAAGTG 151 TCCAGGCTGT GTCTGCTCTT ATTTTAGTGA CAGATGTTGC TCCTGACAGA 201 AGCTATTCTT CAGGAAACAT CACATCCAAT ATGGTAAATC CATCAAACAG 251 GAGCTAAGAA ACAGGAATGA GATGGGCACT TGCCCAAGGA AAAATGCCAG 301 GAGAGCAAAT AATGATGAAA AATAAACTTT TCCCTTTGTT TTTAATTTCA 351 GGAAAAAATG ATGAGGACCA AAATCAATGA ATAAGGAAAA CAGCTCAGAA 401 AAAAGATGTT TCCAAATTGG TAATTAAGTA TTTGTTCCTT GGGAAGAGAC 451 CTCCATGTGA GCTTGATGGG AAAATGGGAA AAACGTCAAA AGCATGATCT 501 GATCAGATCC CAAAGTGGAT TATTATTTTA AAAACCAGAT GGCATCACTC 551 TGGGGAGGCA AGTTCAGGAA GGTCATGTTA GCAAAGGACA TAACAATAAC 601 AGCAAAATCA AAATTCCGCA AATGCAGGAG GAAAATGGGG ACTGGGAAAG 651 CTTTCATAAC AGTGATTAGG CAGTTGACCA TGTTCGCAAC ACCTCCCCGT 701 CTATACCAGG GAACACAAAA ATTGACTGGG CTAAGCCTGG ACTTTCAAGG 751 GAAATATGAA AAACTGAGAG CAAAACAAAA GACATGGTTA AAAGGCAACC 801 AGAACATTGT GAGCCTTCAA AGCAGCAGTG CCCCTCAGCA GGGACCCTGA 851 GGCATTTGCC TTTAGGAAGG CCAGTTTTCT TAAGGAATCT TAAGAAACTC 901 TTGAAAGATC ATGAATTTTA ACCATTTTAA GTATAAAACA AATATGCGAT 951 GCATAATCAG TTTAGACATG GGTCCCAATT TTATAAAGTC AGGCATACAA 1001 GGATAACGTG TCCCAGCTCC GGATAGGTCA GAAATCATTA GAAATCACTG 1051 TGTCCCCATC CTAACTTTTT CAGAATGATC TGTCATAGCC CTCACACACA 1101 GGCCCGATGT GTCTGACCTA CAACCACATC TACAACCCAA GTGCCTCAAC 1151 CATTGTTAAC GTGTCATCTC AGTAGGTCCC ATTACAAATG CCACCTCCCC 1201 TGTGCAGCCC ATCCCGCTCC ACAGGAAGTC TCCCCACTCT AGACTTCTGC 1251 ATCACGATGT TACAGCCAGA AGCTCCGTGA GGGTGAGGGT CTGTGTCTTA

FIG.3A

1301 CACCTACCTG TATGCTCTAC ACCTGAGCTC ACTGCAACCT CTGCCTCCCA 1351 GGTTCAAGCA ATTCTCCTGT CTCAGCCTCC CGCGTAGCTG GGACTACAGG 1401 CGCACGCCCG GCTAATTTTT GTATTGTTAG TAGAGATGGG GTTTCACCAT 1451 ATTAGCCCGG CTGGTCTTGA ACTCCTGACC TCAGGTGATC CACCCACCTC 1501 AGCCTCCTAA AGTGCTGGGA TTACAGGCAT GAGTCACCGC GCCCGGCCAA 1551 GGGTCAGTGT TTAATAAGGA ATAACTTGAA TGGTTTACTA AACCAACAGG 1601 GAAACAGACA AAAGCTGTGA TAATTTCAGG GATTCTTGGG ATGGGGAATG 1651 GTGCCATGAG CTGCCTGCCT AGTCCCAGAC CACTGGTCCT CATCACTTTC 1701 TTCCCTCATC CTCATTTTCA GGCTAAGTTA CCATTTTATT CACCATGCTT 1751 TTGTGGTAAG CCTCCACATC GTTACTGAAA TAAGAGTATA CATAAACTAG 1801 TTCCATTTGG GGCCATCTGT GTGTGTGTAT AGGGGAGGAG GGCATACCCC 1851 AGAGACTCCT TGAAGCCCCC GGCAGAGGTT TCCTCTCCAG CTGGGGGAGC 1901 CCTGCAAGCA CCCGGGGTCC TGGGTGTCCT GAGCAACCTG CCAGCCCGTG 1951 CCACTGGTTG TITTGTTATC ACTCTCTAGG GACCTGTTGC TTTCTATTTC 2001 TGTGTGACTC GTTCATTCAT CCAGGCATTC ATTGACAATT TATTGAGTAC 2051 TTATATCTGC CAGACACCAG AGACAAAATG GTGAGCAAAG CAGTCACTGC 2101 CCTACCTTCG TGGAGGTGAC AGTTTCTCAT GGAAGACGTG CAGAAGAAAA 2151 TTAATAGCCA GCCAACTTAA ACCCAGTGCT GAAAGAAAGG AAATAAACAC 2201 CATCTTGAAG AATTGTGCGC AGCATCCCTT AACAAGGCCA CCTCCCTAGC 2251 GCCCCCTGCT GCCTCCATCG TGCCCGGAGG CCCCCAAGCC CGAGTCTTCC 2301 AAGCCTCCTC CTCCATCAGT CACAGCGCTG CAGCTGGCCT GCCTCGCTTC 2351 CCGTGAATCG TCCTGGTGCA TCTGAGCTGG AGACTCCTTG GCTCCAGGCT 2401 CCAGAAAGGA AATGGAGAGG GAAACTAGTC TAACGGAGAA TCTGGAGGGG 2451 ACAGTGTTTC CTCAGAGGGA AAGGGGCCTC CACGTCCAGG AGAATTCCAG 2501 GAGGTGGGGA CTGCAGGGAG TGGGGACGCT GGGGCTGAGC GGGTGCTGAA 2551 AGGCAGGAAG GTGAAAAGGG CAAGGCTGAA GCTGCCCAGA TGTTCAGTGT 2601 TGTTCACGGG GCTGGGAGTT TTCCGTTGCT TCCTGTGAGC CTTTTTATCT

FIG.3B

2651 TTTCTCTGCT TGGAGGAGAA GAAGTCTATT TCATGAAGGG ATGCAGTTTC 2701 ATAAAGTCAG CTGTTAAAAT TCCAGGGTGT GCATGGGTTT TCCTTCACGA 2751 AGGCCTTTAT TTAATGGGAA TATAGGAAGC GAGCTCATTT CCTAGGCCGT 2801 TAATTCACGG AAGAAGTGAC TGGAGTCTTT TCTTTCATGT CTTCTGGGCA 2851 ACTACTCAGC CCTGTGGTGG ACTTGGCTTA TGCAAGACGG TCGAAAACCT 2901 TGGAATCAGG AGACTCGGTT TTCTTTCTGG TTCTGCCATT GGTTGGCTGT 2951 GCGACCGTGG GCAAGTGTCT CTCCTTCCCT GGGCCATAGT CTTCTCTGCT 3001 ATAAAGACCC TTGCAGCTCT CGTGTTCTGT GAACACTTCC CTGTGATTCT 3051 CTGTGAGGGG GGATGTTGAG AGGGGAAGGA GGCAGAGCTG GAGCAGCTGA 3101 GCCACAGGGG AGGTGGAGGG GGACAGGAAG GCAGGCAGAA GCTGGGTGCT 3151 CCATCAGTCC TCACTGATCA CGTCAGACTC CAGGACCGAG AGCCACAATG 3201 CTTCAGGAAA GCTCAATGAA CCCAACAGCC ACATTTTCCT TCCCTAAGCA 3251 TAGACAATGG CATTTGCCAA TAACCAAAAA GAATGCAGAG ACTAACTGGT 3301 GGTAGCTTTT GCCTGGCATT CAAAAACTGG GCCAGAGCAA GTGGAAAAATG 3351 CCAGAGATTG TTAAACTTTT CACCCTGACC AGCACCCCAC GCAGCTCAGC 3401 AGTGACTGCT GACAGCACGG AGTGACCTGC AGCGCAGGGG AGGAGAAGAA 3451 AAAGAGAGGG ATAGTGTATG AGCAAGAAAG ACAGATTCAT TCAAGGGCAG 3501 TGGGAATTGA CCACAGGGAT TATAGTCCAC GTGATCCTGG GTTCTAGGAG 3551 GCAGGGCTAT ATTGTGGGGG GAAAAAATCA GTTCAAGGGA AGTCGGGAGA 3601 CCTGATTTCT AATACTATAT TTTTCCTTTA CAAGCTGAGT AATTCTGAGC 3651 AAGTCACAAG GTAGTAACTG AGGCTGTAAG ATTACTTAGT TTCTCCTTAT 3701 TAGGAACTCT TTTTCTCTGT GGAGTTAGCA GCACAAGGGC AATCCCGTTT 3751 CTTTTAACAG GAAGAAAACA TTCCTAAGAG TAAAGCCAAA CAGATTCAAG 3801 CCTAGGTCTT GCTGACTATA TGATTGGTTT TTTGAAAAAT CATTTCAGCG 3851 ATGTTTACTA TCTGATTCAG AAAATGAGAC TAGTACCCTT TGGTCAGCTG 3901 TAAACAAACA CCCATTTGTA AATGTCTCAA GTTCAGGCTT AACTGCAGAA 3951 CCAATCAAAT AAGAATAGAA TCTTTAGAGC AAACTGTGTT TCTCCACTCT

FIG.3C

4001 GGAGGTGAGT CTGCCAGGGC AGTTTGGAAA TATTTACTTC ACAAGTATTG 4051 ACACTGTTGT TGGTATTAAC AACATAAAGT TGCTCAAAGG CAATCATTAT 4101 TTCAAGTGGC TTAAAGTTAC TTCTGACAGT TTTGGTATAT TTATTGGCTA 4151 TTGCCATTTG CTTTTTGTTT TTTCTCTTTG GGTTTATTAA TGTAAAGCAG 4201 GGATTATTAA CCTACAGTCC AGAAAGCCTG TGAATTTGAA TGAGGAAAAA 4251 ATTACATTTT TGTTTTTACC ACCTTCTAAC TAAATTTAAC ATTTTATTCC 4301 ATTGCGAATA GAGCCATAAA CTCAAAGTGG TAATAACAGT ACCTGTGATT 4351 TTGTCATTAC CAATAGAAAT CACAGACATT TTATACTATA TTACAGTTGT 4401 TGCAGATACG TTGTAAGTGA AATATTTATA CTCAAAACTA CTTTGAAATT 4451 AGACCTCCTG CTGGATCTTG TTTTTAACAT ATTAATAAAA CATGTTTAAA 4501 ATTTGATAT TTTGATAATC ATATTTCATT ATCATTTGTT TCCTTTGTAA 4551 TCTATATTTT ATATATTTGA AAACATCTTT CTGAGAAGAG TTCCCCAGAT 4601 TTCACCAATG AGGTTCTTGG CATGCACACA CACAGAGTAA GAACTGATTT 4651 AGAGGCTAAC ATTGACATTG GTGCCTGAGA TGCAAGACTG AAATTAGAAA 4701 GTTCTCCCAA AGATACACAG TTGTTTTAAA GCTAGGGGTG AGGGGGGAAA 4751 TCTGCCGCTT CTATAGGAAT GCTCTCCCTG GAGCCTGGTA GGGTGCTGTC 4801 CTTGTGTTCT GGCTGGCTGT TATTTTTCTC TGTCCCTGCT ACGTCTTAAA 4851 GGACTTGTTT GGATCTCCAG TTCCTAGCAT AGTGCCTGGC ACAGTGCAGG 4901 TTCTCAATGA GTTTGCAGAG TGAATGGAAA TATAAACTAG AAATATATCC 4951 TTGTTGAAAT CAGCACACCA GTAGTCCTGG TGTAAGTGTG TGTACGTGTG 5001 TGTGTGTGT TGTGTGTGT TGTAAAACCA GGTGGAGATA TAGGAACTAT 5051 TATTGGGGTA TGGGTGCATA AATTGGGATG TTCTTTTTAA AAAGAAACTC 5101 CAAACAGACT TCTGGAAGGT TATTTTCTAA GAATCTTGCT GGCAGCGTGA 5151 AGGCAACCCC CCTGTGCACA GCCCCACCCA GCCTCACGTG GCCACCTCTG 5201 TCTTCCCCCA TGAAGGGCTG GCTCCCCAGT ATATATAAAC CTCTCTGGAG 5251 CTCGGGCATG AGCCAGCAAG GCCACCCATC CAGGCACCTC TCAGCACAGC 5300

FIG.3D

(exon #2) TT TCT ACG TGG AAT TTG GAC

661 ACT TTG GCCTTC CAG GAA CTG AAG TCC GAG CTA ACT GAA GTT CCT GCTTCC CGA ATT TTG 720 721 AAG GAG AGC CCA TCT GGCTAT CTC AGG AGT GGA GAG GGA GAC ACCG

FIG.3E

tggctctgccaagcttccgcatgatcattgtctgtgtttggaagattatggattaagtggtgcttcgtttt ctttctgaatttaccag

(exon #3) GA TGT GGA GAA CTA 780

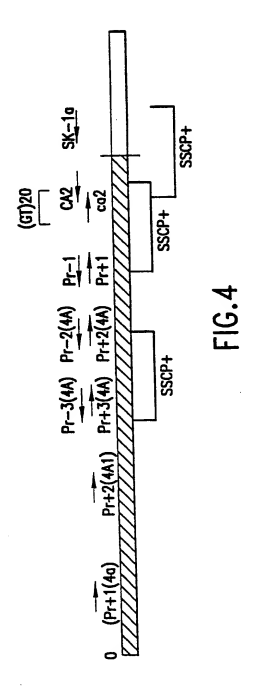
940 900 960 1020 1140 1260 1320 1380 1440 46A 46A 66CT 66CT 77T 77C 66CT 66CT 66CT ACA CAG GAC CTG GGCT GGCT CGT TAC CTG ACA CCC TTT CCT GCT GGT GGT GTC ATC TAC ATC CTG ACC CAG CAG GCC GCC ACA ATT ATT ACG CCC CGC CGC CAC TTC ACT ACC ACC ACC ACC CTC AAG GTT TAT TAT CAA CAA ACC ACC GTC CCC CCC GAT CTC GGG GGT GGC GGC GGC ACC GGA GGG GGG GGG GAG TCT TCT TCT TCT GAA GAA AAG 6TA 4TG 6TT 1CG 1CG ACC ATC ATC ATC ATC 166 ACA AAT CCCG GTC GTC GAC GAC 611 616 646 646 676 646 660 670 640 670 781 841 901 901 1021 1081 1141 1201 1321 1381 1381 1441

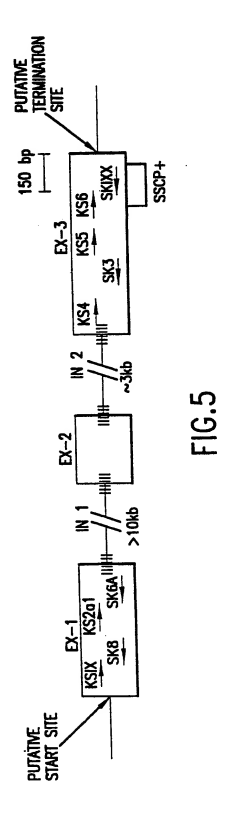
TCC 1560 flanking region) TGA AAA GCC . E 1620 1680 1740 1800 1860 1920 1980 2040 2100 2160 CTG 17AA 17AG 6AC CTG CTG 17C AAG AGCTT AGG GGG AGT CTG AGT AAG AGT TGC CCCT TGT GGG CCA AGT AAA AGA AGA AGA AGA GCT 16C AAT AAT 17G CAC GTTCA ATG CTA AAA ACC CAT CCT TTC 646 667 144 671 671 740 671 740 740 AAG CCA ATC TTA ATG GGC GAA TCT AGG CAG GGC ACC CCT TTT TAC ATA ATC 766 CAG GTC GTC TGT TGT ACC CAA ACC CTG GAG AGG CAA ATT CTT CTT CTT AAG AGG AGG AGG AGG AGG AGG 1561 1621 1681 1741 1861 1981 1981 2041 2101

FIG.3F

2581 ATG TTA GCC AGG ATG GTC TCG ATC TCC TGAC CTT GTC ATC CAC CCA CCT TGG CCT CCC AAA 2640 2641 GTG CTG GGA TTA CAG GCG TGA GCC ACC ACGC CCA GCC CCT CCA CTT CAG TTT TTA TCT GTC 2700 CAG GCA CCT GCC ACG CCT GGC TAA TTT TTT TTT CCA GTG AAG ATG GGT TTC ACC 2580 2701 ATC AGG GGT ATG AAT TTT ATA AGC CAC ACC TCA GGT GGA GAA AGC TTG ATG CAT AGC TTG 2760 2401 GAT GGG GTC TCG CTC TGT CAC CCA GGC TGG AGT GCA GTG GCA CGA TCT CGG CTC ACT GCA 2460 AGT TCC GCC TCC CAG GTT CAC ACC ATT CTC CTG CCT CAG CCT CCC AAG TAG CTG GGA CTA 2520 2281 CTG GTG CTC TCG GCA CTA CCG GAG CAA TCT TTC CAT CTC TCC CCT GAA CCC ACC CTC TAT 2340 TCC CAC CTC CAT CTC 2280 TTA AAG GTG TGT GTG TGT GTG TGT GTG TAA CTG AGA GGC TTG TGC CTG GTT TTG 2220 TCA CCC TAA CTC CAC TTC AGT TTG CTT TTG ATT TTT TTT TTT TTT TTT TTT TGA AGG TGC TGC CCA GGA TGA CGC CAA GCA AAT AGC GCA TCC ACA CTT ATT CTA TAC TGT 2776 2761 AGT 2341 2161

FIG.3G





23 23 23 23 23	119 97 120 119 120	178 131 178 177 177
-TGAVVYSGS LYFQGAESRT VIRYELNTET VKAEKEIPGA GYHGQFPYSW GGYTDIDLAV 59	DEAGLWVIYS TDEAKGAIVL SKLNPENLEL EQTWETNIRK QSVANAFIIC GTLYTVSSYT 119 DESGLWAVYA TNQNAGNIVV SRLDPVSLQT LQTWNTSYPK RXPGXAFIIC GTCYVTNGY- 97 DENGLWAVYA TNQNAGNIVI SKLDPVSLQI LQTWNTSYPK RSAGEAFIIC GTLYVTNGYS 120 DEKGLWVIFT TEKSAGKIVV GKVNVATFTV DNIWITTQNK SDASNAFMIC GVLYVTRSLG 119 DE.GLWY. TAG.IV. SKL.PL QTW.TKAFIIC GTLYVTY. 120	MVTYDIKLSHFPC QTLYNVTLF LLHYDIAL-
GYHGQFPYSW GYNNMYHYAW GYNNMYHYAW LFNNRFSYAG GYNNYAW	QSVANAFIIC RXPGXAFIIC RSAGEAFIIC SDASNAFMIC	SADATVNFAY DTGTGISKTL TIPFKNRYKY SSMIDYNPLE KKLFAWDNLN MVTYDIKLS SGGTKVHYAY QTNASTYEYIDI-PFQ NKLXPHFPC GG-TKVHYAY QTNASTYEYI DIPFQNKYSH ISMLDYNPKD RALYAWNNGH QTLYNVTLF PKMEEVFYMF DTKTGKEGHL SIMMEKMAEK VHSLSYNSND RKLYMFSEGY LLHYDIAL- NYAY TY.
VKAEKE IPGA ILKTRSLDYA ILKTRSLDYA VYQKKPLLNA LA	EQTWETNIRK LQTWNTSYPK LQTWNTSYPK DNIWITTQNK	SSMIDYNPLEIDI-PFQ ISMLDYNPKD VHSLSYNSND
VIRYELNTETRFDLKTET IIRFDLKTET MCRASL-TSGRL.TET	SKLNPENLEL SRLDPVSLQT SKLDPVSLQI GKVNVATFTV SKL.PL.:	TIPFKNRYKY YEY DIPFQNKYSH SIMMEKMAEK
LYFQGAESRT IYFNKFQSHI LYYNCFNSHD YS	TDEAKGAIVL TNQNAGNIVV TNQNAGNIVI TEKSAGKIVV TAG.IV:	DTGTG1SKTL QTNAST QTNASTYEYI DTKTGKEGHL . T
-TGAVVYSGS GTGQVVYNGS GAGVVVHNNN .G.VV	DEAGLWVIYS DESGLWAVYA DENGLWAVYA DEKGLWVIFT DE.GLWY.	SADATVNFAY SGGTKVHYAY GG-TKVHYAY PKMEEVFYMF
TIGR ymOBhl2.rl 1B426bAMZ ranofm Consensus	TIGR ym08h12.rl 1B426bAMZ ranofm Consensus	TIGR ymO8h12.rl 1B426bAMZ ranofm Consensus

FIG.6

TTG GCC TTC CAG GAA CTG AAG TCC GAG CTA ACT GAA GTT CCT GCT TCC CGA ATT TTG 720 540 TTG GAA AGC AGC CAG CAG GTA GCA AGG CTG AGA AGG GGC CAG TGT CCC CAG ACC 600 ACG TGG AAT TTG GAC 660 GGA GAA CTA 780 420 480 240 300 360 CCT GAG ATG CCA GCT GTC CAG CTG CTT CTG GCC TGC CTG GTG TGG 120 CGA TGC CAG 180 GTG TGG ATG CGA GAC CCC AAG CCC ACC TAC CCC TAC ACC CAG GAG ACC ACG TGG AGA ATC GTT TGG GTA GGA GAG CCT CTC ACG CTG AGA ACA GCA GAA ACA ATT ACT GGC AAG TAT GGT 421 GAC CAG CTG GAA ACC CAA ACC AGA GAG TTG GAG ACT GCC TAC AGC AAC CTC CTA GAC **A**66 CCC AAT GAA TCC AGC TGC CCA GAG CAG AGC CAG GCC ATG ACC AAA GCT CGA CTC AGC TCC CTG GAG AGC CTC CAC CAA TTG ACC TTG GAC CAG GCT GCC AGG CCC CAG GAG ACC CAG GAG GGG CTG CAG AGG GAG CTG GGC ACC CTG AGG CGG GAG CGG 1 AGA GCT TTC CAG AGG AAG CCT CAC CAA GCC TCT GCA ATG AGG TTC TTC TGT GCA CGT TGC TCA GTC ATC CAT AAC TTA CAG AGA GAC AGC AGC CAA CGC TTA GAC CTG GAG GCC GAG AAT CTG GCC AAG GAG AGC CCA TCT GGC TAT CTC AGG AGT GGA GAG GGA GAC ACC GGA TGT ည္ဟ GAT GTG GGG GCC AGG ACA GCT CAG CTC AGG AAG GCC AAT GAC CAG AGT CGA GAC ACT GCT CGG GCT GTG CCA CCA GGC TCC AGA GAA GTT TCT 481 AAG TCA GTT CTG GAG GAA GAG AAG AAG CGA CTA AGG CAA GAA AAT TTC AGT GTG GCC AGT 999 E 181 TAT ACC AGC 721 361 661 601

FIG.7A

1548 GCC TTC ATC ATC TGT GGC ACC TTG TAC ACC GTC AGC TAC ACC TCA GCA GAT GCT ACC 1380 1381 GTC AAC TTT GCT TAT GAC ACA GGC ACA GGT ATC AGC AAG ACC CTG ACC ATC CCA TTC AAG 1440 1441 AAC CGC TAT AAG TAC AGC AGC ATG ATT GAC TAC AAC CCC CTG GAG AAG AAG CTC TTT GCC 1500 GTG TAC TCG GGG AGC CTC TAT TTC CAG GGC GCT GAG TCC AGA ACT GTC ATA AGA TAT GAG 1080 CTG AAT ACC GAG ACA GTG AAG GCT GAG AAG GAA ATC CCT GGA GCT GGC TAC CAC GGA CAG 1140 1141 TTC CCG TAT TCT TGG GGT GGC TAC ACG GAC ATT GAC TTG GCT GTG GAT GAA GCA GGC CTC 1200 1201 TGG GTC ATT TAC AGC ACC GAT GAG GCC AAA GGT GCC ATT GTC CTC TCC AAA CTG AAC CCA 1260 GAG AAT CTG GAA CTC GAA CAA ACC TGG GAG ACA AAC ATC CGT AAG CAG TCA GTC GCC AAT 1320 CAG GGC TAC CCT TCT AAG GTT CAC ATA CTG CCT AGG CCA CTG GAA AGC ACG GGT GCT GTG 1020 960 901 GAC ACA GTT GGC ACG GAT GTC CGC CAG GTT TTT GAG TAT GAC CTC ATC AGC CAG TTT ATG TTG AAC ATG GTC ACT TAT GAC ATC AAG CTC TCC AAG ATG 1501 TGG GAC AAC 1321

FIG. 7E

Met Arg Phe Phe Cys Ala Arg Cys 20

221 Thr Leu Ala Phe Gln Glu Leu Lys Ser Glu Leu Thr Glu Val Pro Ala Ser Arg Ile Leu 240 241 Lys Glu Ser Pro Ser Gly Tyr Leu Arg Ser Gly Glu Gly Asp Thr Gly Cys Gly Glu Leu 260 261 Val Trp Val Gly Glu Pro Leu Thr Leu Arg Thr Ala Glu Thr Ile Thr Gly Lys Tyr Gly 280 281 Val Irp Met Arg Asp Pro Lys Pro Thr Tyr Pro Tyr Thr Gln Glu Thr Thr Trp Arg Ile 300 181 Arg Leu Glu Ser Ser Ser Gln Glu Val Ala Arg Leu Arg Arg Gly Gln Cys Pro gln Thr 200 121 Arg Pro Gln Glu Thr Gln Glu Gly Leu Gln Arg Glu Leu Gly Thr Leu Arg Arg Glu Arg 140 201 Arg Asp Thr Ala Arg Ala Val Pro Pro Gly Ser Arg Glu Val Ser Thr Trp Asn Leu Asp 220 101 Lys Ala Arg Leu Ser Ser Leu Glu Ser Leu Leu His Gln Leu Thr Leu Asp Gln Ala Ala 120 141 Asp Gln Leu Glu Thr Gln Thr Arg Glu Leu Glu Thr Ala Tyr Ser Asn Leu Leu Arg Asp 160 161 Lys Ser Val Leu Glu Glu Glu Lys Lys Arg Leu Arg Gln Glu Asn Glu Asn Leu Ala Arg 180 81 Ser Val Ile His Asn Leu Gln Arg Asp Ser Ser Thr Gln Arg Leu Asp Leu Glu Ala Thr 100 61 Tyr Thr Phe Ser Val Ala Ser Pro Asn Glu Ser Ser Cys Pro Glu Gln Ser Gln Ala Met 80 41 Asp Val Gly Ala Arg Thr Ala Gln Leu Arg Lys Ala Asn Asp Gln Ser Gly Arg Cys Gln 60 21 Cys Ser Phe Gly Pro Glu Met Pro Ala Val Gln Leu Leu Leu Leu Ala Cys Leu Val Trp 40 FIG.8A

500 Ala Phe Ile Ile Cys Gly Thr Leu Tyr Thr Val Ser Ser Tyr Thr Ser Ala Asp Ala Thr 460 461 Val Asn Phe Ala Tyr Asp Thr Gly Thr Gly Ile Ser Lys Thr Leu Thr Ile Pro Phe Lys 480 381 Phe Pro Tyr Ser Trp Gly Gly Tyr Thr Asp Ile Asp Leu Ala Val Asp Glu Ala Gly Leu 400 401 Trp Val Ile Tyr Ser Thr Asp Glu Ala Lys Gly Ala Ile Val Leu Ser Lys Leu Asn Pro 420 361 Leu Asn Thr Glu Thr Val Lys Ala Glu Lys Glu Ile Pro Gly Ala Gly Tyr His Gly Gln 380 340 341 Val Tyr Ser Gly Ser Leu Tyr Phe Gln Gly Ala Glu Ser Arg Thr Val Ile Arg Tyr Glu 360 320 Asn Arg Tyr Lys Tyr Ser Ser Met Ile Asp Tyr Asn Pro Leu Glu Lys Lys Leu Phe Ala Glu Asn Leu Glu Leu Glu Gln Thr Trp Glu Thr Asn Ile Arg Lys Gln Ser Val Ala Asn 301 Asp Thr Val Gly Thr Asp Val Arg Gln Val Phe Glu Tyr Asp Leu Ile Ser Gln Phe Met Gln Gly Tyr Pro Ser Lys Val His Ile Leu Pro Arg Pro Leu Glu Ser Thr Gly Ala Val 501 Trp Asp Asn Leu Asn Met Val Thr Tyr Asp Ile Lys Leu Ser Lys Met

FIG.8B

Gel Shift Assays

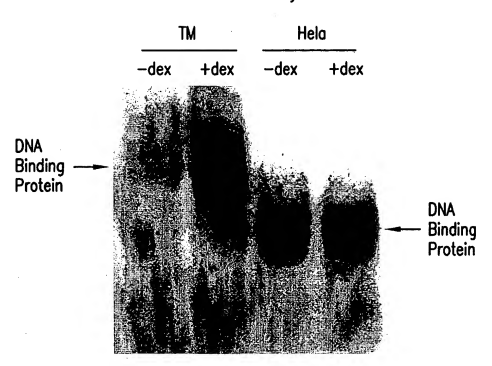


FIG.9A

Gel Shift Assays

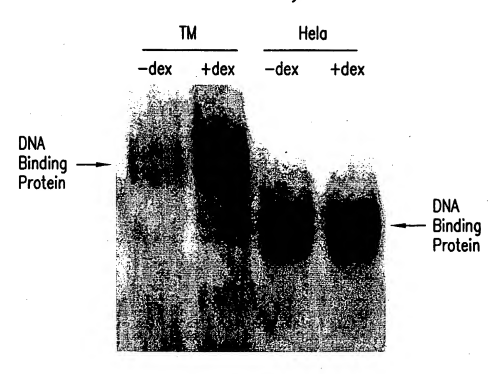


FIG.9B

SEQUENCE LISTING

- (1) GENERAL INFORMATION
- (i) APPLICANT: Nguyen, Thai D. Polansky, Jon R. Chen, Pu Chen, Hua
- (ii) TITLE OF THE INVENTION: NUCLEIC ACIDS, KITS, AND METHODS FOR THE DIAGNOSIS, PROGNOSIS AND TREATMENT OF GLAUCOMA AND RELATED DISORDERS
 - (iii) NUMBER OF SEQUENCES: 38
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Howrey & Simon
 - (B) STREET: 1299 Pennsylvania Avenue, N.W.
 - (C) CITY: Washington
 - (D) STATE: DC
 - (E) COUNTRY: USA
 - (F) ZIP: 20004-2402
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Diskette
 - (B) COMPUTER: IBM Compatible

 - (C) OPERATING SYSTEM: DOS
 (D) SOFTWARE: FastSEQ for Windows Version 2.0
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:

 - (vii) PRIOR APPLICATION DATA:
 (A) APPLICATION NUMBER: 09/227,881
 (B) FILING DATE: 11-JAN-1999
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME:
 - (B) REGISTRATION NUMBER:
 - (C) REFERENCE/DOCKET NUMBER: 07425-0051
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 202 783-0800 (B) TELEFAX: 202 383-6610 (C) TELEX:

 - (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5300 base pairs

 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

60 TCCTATAAAC TGTATAGCCT CCATTCGGAT GTATGTCTTT GGCAGGATGA TAAAGAATCA 120

CC3 3 C3 3 CC3	GTATCCACGT	TAGCCAAGTG	TCCAGGCTGT	GTCTGCTCTT	ATTTTAGTGA	180
CARGORAGOR	TCCTGACAGA	VCCury LunCum	CAGGAAACAT	CACATCCAAT	ATGGTAAATC	240
CAGATGTTGC	GAGCTAAGAA	ACACCAATCA	CATCCCCACT	TGCCCAAGGA	AAAATGCCAG	300
CATCAAACAG	AATGATGAAA	ACAGGAAIGA	TO COMPTONI	ጥጥጥ ልጥጥጥCA	GGAAAAAATG	360
GAGAGCAAA'I'	AATGATGAAA	AMIMAMCIII	CACCHIGIT	A A A A C A TO COTT	TCCAAATTGG	420
ATGAGGACCA	AAATCAATGA	ATAAGGAAAA	CAGCICAGAA	MAMAGAIGII	AAAATCCCAA	480
TAATTAAGTA	TTTGTTCCTT	GGGAAGAGAC	CTCCATGTGA	GCTTGATGGG	AAAAIGGGAA	540
AAACGTCAAA	AGCATGATCT	GATCAGATCC	CAAAGTGGAT	TATTATTTTA	MANACCAGA:	600
GGCATCACTC	TGGGGAGGCA	AGTTCAGGAA	GGTCATGTTA	GCAAAGGACA	TAACAATAAC	
AGCAAAATCA	AAATTCCGCA	AATGCAGGAG	GAAAATGGGG	ACTGGGAAAG	CTTTCATAAC	660
አርመር አመጥአርር	CACTTGACCA	TGTTCGCAAC	ACCTCCCCGT	CTATACCAGG	GAACACAAAA	720
AMMCACTICAG	CTAAGCCTGG	ACTTTCAAGG	GAAATATGAA	AAACTGAGAG	CAAAACAAAA	780
CXCXTCCTTX	AAAGGCAACC	AGAACATTGT	GAGCCTTCAA	AGCAGCAGTG	CCCCTCAGCA	840
CCCACCCTCA	GGCATTTGCC	TTTAGGAAGG	CCAGTTTTCT	TAAGGAATCT	TAAGAAACTC	900
መመር ል ል ል ር ል ጥር	ልጥር ልልጥጥጥTA	ACCATTTTAA	GTATAAAACA	AATATGCGAT	GCATAATCAG	960
TOTAL CACATIC	CCTCCCA ATT	TTATAAAGTC	AGGCATACAA	GGATAACGTG	TCCCAGCTCC	1020
CCAMACCATCA	GAAATCATTA	GAAATCACTG	TGTCCCCATC	CTAACTTTTT	CAGAATGATC	1080
MCMCAMACCC	CTCACACACA	GGCCCGATGT	GTCTGACCTA	CAACCACATC	TACAACCCAA	1140
TGTCATAGCC	CATTGTTAAC	CTCTCATCTC	AGTAGGTCCC	ATTACAAATG	CCACCTCCCC	1200
GTGCCTCAAC	ATCCCGCTCC	ACACCAACTC	TO COCO CTOT	AGACTTCTGC	ATCACGATGT	1260
TGTGCAGCCC	AGCTCCGTGA	COCTORCECCO	COCCACICI	CACCTACCTG	TATGCTCTAC	1320
TACAGCCAGA	AGCTCCGTGA	GGGTGAGGGT	CIGICILIA	AMMONOCHOM	CTCAGCCTCC	1380
ACCTGAGCTC	ACTGCAACCT	CTGCCTCCCA	GGTTCAAGCA	ATTOTOGIGE	MACAGATGGG	1440
CGCGTAGCTG	GGACTACAGG	CGCACGCCCG	GCTAATTTT	GTATIGTTAG	TAGAGATGGG	1500
GTTTCACCAT	ATTAGCCCGG	CTGGTCTTGA	ACTCCTGACC	TCAGGTGATC	CACCCACCTC	1560
AGCCTCCTAA	AGTGCTGGGA	TTACAGGCAT	GAGTCACCGC	GCCCGGCCAA	GGGTCAGTGT	
መጠአ አመአ አርር አ	ል ጥ ል ል ር ጥጥር ል ል	ጥርርጥጥጥልርጥል	AACCAACAGG	GAAACAGACA	AAAGCTGTGA	1620
ጥል አመጥጥር ልርር	CATTCTTGGG	ATGGGGAATG	GTGCCATGAG	CTGCCTGCCT	AGTCCCAGAC	1680
CACTGGTCCT	CATCACTTTC	TTCCCTCATC	CTCATTTTCA	GGCTAAGTTA	CCATTTATT	1740
CACCAMCCMI	THE THE CONTRACT	CCTCCACATC	GTTACTGAAA	TAAGAGTATA	CATAAACTAG	1800
መጥረር አጥጥጥርር	CCCCATCTGT	GTGTGTGTAT	AGGGGAGGAG	GGCATACCCC	AGAGACTCCT	1860
MC A ACCCCCC	CCCACAGGGTT	TCCTCTCCAG	CTGGGGGAGC	CCTGCAAGCA	CCCGGGGGTCC	1920
TO CONTRACTOR	CAGCAACCTG	CCAGCCCGTG	CCACTGGTTG	TTTTGTTATC	ACTCTCTAGG	1980
CACCOCOTOC		TGTGTGACTC	GTTCATTCAT	CCAGGCATTC	ATTGACAATT	2040
GACCIGIIGC	, ILICIALITE	CAGACACCAG	AGACAAAATG	GTGAGCAAAG	CAGTCACTGC	2100
TATTGAGTAC	, TIMIMICIGO	A CHAMACACOTTO	CCAACACGTG	CAGAAGAAAA	TTAATAGCCA	2160
CCTACCTTCG	TGGAGGTGAC	AGIIICICAI	מרבונים מרבונים ביים ביים ביים ביים ביים ביים ביים ב	CATCTTGAAG	AATTGTGCGC	2220
GCCAACTTAA	ACCCAGTGCT	COMPONIANCE	CCCCCCCTCCT	CCCTCCATC	TGCCCGGAGG	2280
AGCATCCCT	· AACAAGGCCA	LOTCCCTAGC	CECCOCIOCI	CACACCCCTC	CAGCTGGCCT	2340
CCCCCAAGC	CGAGTCTTCC	AAGCCTCCTC	. CICCAICAGI	ACACHGCGCTC	COTCCAGGCT	2400
GCCTCGCTTC	CCGTGAATCG	TCCTGGTGCA	TCTGAGCTGG	MCMCCACCCC	GCTCCAGGCT	2460
CCAGAAAGG	A AATGGAGAGG	GAAACTAGTC	TAACGGAGAA	TCTGGAGGGG	ACAGTGTTTC	2520
CTCAGAGGG	A AAGGGGCCTC	CACGTCCAGG	AGAATTCCAG	GAGGTGGGG	CTGCAGGGAG	2580
TGGGGACGC'	r ggggctgagc	GGGTGCTGAA	AGGCAGGAAG	GIGAAAAGGG	CAAGGCTGAA	2640
GCTGCCCAG	A TGTTCAGTGT	TGTTCACGGG	GCTGGGAGT"	TTCCGTTGCT	TCCTGTGAGC	2700
CTTTTTATC	r TTTCTCTGCT	' TGGAGGAGAA	GAAGTCTATT	TCATGAAGG	ATGCAGTTTC	
አጥአ አአርጥሮ እር	2 ሮሞርሞሞልልልልባ	TOTAGGGTG1	' GCATGGGTTI	TCCTTCACGA	AGGCCTTTAT	2760
ጥጥል አጥርርርርል :	TATACCAACC	GAGCTCATT	CCTAGGCCG	TAATTCACGO	AAGAAGTGAC	2820
ጥርር እርጥርጥጥ	ው መርመመምር <u>አመር</u> ባ	* CTTCTGGGC <i>I</i>	A ACTACTCAGO	CCTGTGGTG	ACTIGGCTTA	2880
TO CARCACO	תרפאאאאררי	THE TENDER TO THE	AGACTCGGTT	P TTCTTTCTG	3 TTCTGCCATT	2940
COMMCCOMO	T CCGACCGTGC	CCAAGTGTC1	r creerrece	r gggccatag	r Cricicitati	3000
አጥአ አልርልሮር	ר התככם בכתכי	r ტვუცუულე	r GAACACTTC	CTGTGATTC'	r CTGTGAGGGG	3060
CCAMOMOCO	C ACCCCAACC	GCAGAGCTO	GAGCAGCTG	A GCCACAGGG	AGGTGGAGGG	3120
GGATGIIGA	C CCACCCACA	COTCCCTCC	T CCATCAGTC	TCACTGATC	A CGTCAGACTC	3180
GGACAGGAA	G GCAGGCAGA	COMCACCAA	CCTCAATGA	CCCAACAGC	CACATTTTCCT	3240
CAGGACCGA	G AGCCACAAT	2 CITCAGGAA	A MANCONANA	A GAATGCAGA	G ACTAACTGGT	3300
TCCCTAAGC	A TAGACAATG	CATTIGCCA	A IMACCAMAM	A CECCANANT	CCACACATTC	3360
GGTAGCTTT	T GCCTGGCAT	r caaaaacig	GCCAGAGCA	A GIGGAAAAI	G CCAGAGATTG	3420
TTAAACTTT	T CACCCTGAC	C AGCACCCCA	C GCAGCTCAG	C AGTGACTGC	T GACAGCACGG	3480
AGTGACCTG	C AGCGCAGGG	g aggagaaga	A AAAGAGAGG	G ATAGTGTAT	G AGCAAGAAAG	3540
ACAGATTCA	T TCAAGGGCA	G TGGGAATTG.	A CCACAGGGA	T TATAGTCCA	C GTGATCCTGG	3600
GTTCTAGGA	G GCAGGGCTA	T ATTGTGGGG	G GAAAAAATC	A GTTCAAGGG	A AGTCGGGAGA	
ഗാനവു സന്നാ	ጥ አአጣአርጥአጥል	տ փափանանական ա	A CAAGCTGAG	T AATTCTGAG	C AAGTCACAAG	3660
CMACMAACT	יכ אככריייכיייא א	ር አጥጥእርጥጥእር	Т ТОРГОТОСТТА	T TAGGAACTC	T TTTTCTCTGT	3720
CCACTTACC	A CCACAACC	ሮ አአጥሮሮሮርጥጥ	т сттттааса	G GAAGAAAAC	A TTCCTAAGAG	3780
ጥል እ አርርር እ እ	ል ሮልፎልሞሞሮልል	G CCTAGGTCT	T GCTGACTAT	A TGATTGGTT	T TTTGAAAAT	3840
ር እ ጥጥጥር እርር	ነር አጥርጥጥጥአርጥ	A TOTGATTCA	G AAAATGAGA	C TAGTACCUI	T TEGTCAGCTG	3900
መአአአሮአአልር	יא כככאייייייים מי	ል ልልጥርጥርጥርል	A GTTCAGGCT	T AACTGCAGA	A CCAATCAAAT	3960
TAMACAMAC	T CCCUTTIGE	C ALACTOTOR	T TOTOCACTO	T GGAGGTGAG	T CTGCCAGGGC	4020
AAGAATAGA	ry iclitaded	C VCVVCIGIGI	C ACACTICTOR	T TGGTATTAA	C AACATAAAGT	4080
AGTTTGGAA	AA TATTTACTI	M MACAWATATI	C TOTAL ACTOR	C TTCTGACAG	T TTTGGTATAT	4140
TGCTCAAAC	G CAATCATTA	T IICMMGIGG	2			
			7)			

PCT/US00/00559 WO 00/42220

(2) INFORMATION FOR SEQ ID NO:2:

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 5304 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

TO THE TAX	ACC AATGTGAAAG 60	
AMOMMMOCOTO AGTITACCIC AGGGCTITTION	TGA TAAAGAATCA 120	
TCCTATAAAC TGTATAGCCT CCATTCGGAT GTATGTCTTT GGCAGGA	CTT ATTTTAGTGA 180	
CCARGACCA GTATCCACGT TAGCCAAGTG TCCAGGCTGT GTCTGCT	240	
CACATGTTCC TCCTGACAGA AGCTATTCTT CAGGAAACAT CACATCC	GGA AAAATGCCAG 300	
CATCA ACAG GAGCTAAGAA ACAGGAATGA GATGGGCACT TGCCCAA	TCA GGAAAAATG 360	
CACACCADAT AATGATGAAA AATAAACTTT TCCCTTTGTT TITAATI		
ATGAGGACCA AAATCAATGA ATAAGGAAAA CAGCTCAGAA AAAAGAT	400	
TAATTAAGTA TTTGTTCCTT GGGAAGAGAC CTCCATGIGA GCTIONI	TTTA AAAACCAGAT 540	
AAACGTCAAA AGCATGATCT GATCAGATCC		
CCCATCACTC TGGGGAGGCA AGTICAGGAA GGICALI		
AGCAAAATCA AAATTCCGCA AATGCAGGAG GAATTCCGCG CMATTACC	700	
AGTGATTAGG CAGTTGACCA TGTTCGCAAC ACCTCCCGI CIACAC	700	
ATTGACTGGG CTAAGCCTGG ACTTTCAAGG CAAGCCT	040	
GACATGGTTA AAAGGCAACC AGAACATTGT GACGAT		
GGGACCCTGA GGCATTTGCC TTTAGGAAGG CCAGTITTCT TAMOUNT		
TTGAAAGATC ATGAATTTTA ACCATTTTAA GTATAAACA ACTATAA	CGTG TCCCAGCTCC 1020	
TTTAGACATG GGTCCCAATT TTATAAAGTC ACCESTED CON ACTU	1000	
GGATAGGTCA GAAATCATTA GAAATCACTO	1140	
TGTCATAGCC CTCACACACA GGCCCGATGT GTCTGCG AMTACA	AATG CCACCTCCCC 1200	
GTGCCTCAAC CATTGTTAAC GIGICATCTC	1000	
TGTGCAGCCC ATCCCGCTCC ACAGGAAGIC ICCCCTTT CACGTT	CCTG TATGCTCTAC 1320	
TACAGCCAGA AGCTCCGTGA GGGTGAGGGT		
ACCTGAGCTC ACTGCAACCT CIGCCICCCA		
CGCGTAGCTG GGACTACAGG CGCACGCCCG		
GTTTCACCAT ATTAGCCCGG CIGGICIIGA	1560	
AGCCTCCTAA AGTGCTGGGA TTACAGGCAI GAGTCAGGC	1600	
TTAATAAGGA ATAACTTGAA IGGITIACTA	- 1600	
TAATTTCAGG GATTCTTGGG ATGGGGGAATG CTCATTTTCA GGCTAA	AGTTA CCATTTTATT 1740	
CACTGGTCCT CATCACTTTC TICCCICATE CTMACTCAAA TAACAC	1000	
CACCATGCTT TTGTGGTAAG CCICCACAIC	ACCCC AGAGACTCCT 1860	
TTCCATTTGG GGCCATCTGT GTGTGTGTAT	AAGCA CCCGGGGTCC 1920	
TGAAGCCCC GGCAGAGGTT ICCICIONA GGACGCCTTC TTTTTG	TTATC ACTCTCTAGG 1980	
TGGGTGTCCT GAGCAACCTG CCAGCCCGTG CCACTGOTTCAT CCAGGG	CATTC ATTGACAATT 2040	
GACCTGTTGC TTTCTATTTC IGIGGGC ACACAAATG GTGAG	CAAAG CAGTCACTGC 2100)
TATTGAGTAC TTATATCTGC CAGACACCAG AGACAAAATG GIGAG	•	

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CCTACCTTCG TGGAGGTGAC AGTTTCTCAT GGAAGACGTG CAGAAGAAAA TTAATAGCCA	2160
CCTACCTTCG TGGAGGTGAC AGTTTCTCAT GGAGGACGTG CATCTTGAAG AATTGTGCGC GCCAACTTAA ACCCAGTGCT GAAAGAAAGA AAATAAACAC CATCTTGAAG AATTGTGCGC GCCAACTTAA ACCCAGTGCT GAAAGAAAGA AAATAAACAC CATCCTTGCT GCCTCCATCG TGCCCGGAGG	2220
GCCAACTTAA ACCCAGTGCT GAAAGAAAGG AAATAAACAC CATCCATCG TGCCCGGAGG AGCATCCCTT AACAAGGCCA CCTCCCTAGC GCCCCCTGGT GCCTCCATCG TGCCCGGAGG AGCATCCCTT AACAAGGCCA CCTCCCTAGC CTCCATCAGT CACAGCGCTG CAGCTGGCCT	2280
AGCATCCCTT AACAAGGCCA CCTCCCTAGC GCCCCTCATCAGT CACAGCGCTG CAGCTGGCCT CCCCCAAGCC CGAGTCTTCC AAGCCTCCTC CTCCATCAGT CACAGCGCTG CAGCTGGCT CCCCCAAGCC CGAGTCTTCC AAGCCTCCATCAGT CACAGCGCTTG GCTCCAGGCT CCCCCAAGCC CGAGTCTTCC AAGCCTCCATCAGT CACAGCTTCTTG GCTCCAGGCT	2340
CCCCCAAGCC CGAGTCTTCC AAGCCTCCTC CTCATCAGT AGACTCCTTG GCTCCAGGCT GCCTCGCTTC CCGTGAATCG TCCTGGTGCA TCTGAGCTG AGACTCCTTG GCTCCAGGCT GCCTCGCTTC CCGTGAATCG TCCTGGTGCA TCTGGAGGA ACAGTGTTTC	2400
GCCTCGCTTC CCGTGAATCG TCCTGGTGCT TAACGGTGAA TCTGGAGGGG ACAGTGTTTC	2460
CCAGAAAGGA AATGGAGAGG GAAACTAGTC TARACTAGAG CAGGTGGGGA CTGCAGGGAG	2520
CTCAGAGGGA AAGGGGCCTC CACGTCCAGG AGGCCAAC GTGAAAGGG CAAGGCTGAA	2580
TGGGGACGCT GGGGCTGAGC GGGTGCTGAA AGTGGCAGTT TTCCGTTGCT TCCTGTGAGC	2640
GCTGCCCAGA TGTTCAGTGT TGTTCACGGG GAACTCTATT TCATGAAGGG ATGCAGTTTC	2700
CTTTTTATCT TTTCTCTGCT TGGAGGAGAA GAARGCCTTTAT	2760
ATAAAGTCAG CTGTTAAAAT TCCAGGGTGT GCTACCCCGT TAATTCACGG AAGAAGTGAC	2820
TTAATGGGAA TATAGGAAGC GAGCICATIA	2880
TGGAGTCTTT TCTTCATGT CTTCTGGGCA ACLORECCOUR TTCTTCTGG TTCTGCCATT	2940
TGCAAGACGG TCGAAAACCT TGGAATCAGG RECOTTCGCT GGGCCATAGT CTTCTCTGCT	3000
GGTTGGCTGT GCGACCGTGG GCAAGIGICI CLACAGTGCC CTGTGATTCT CTGTGAGGGG	3060
ATAAAGACCC TTGCAGCTCT CGTGTTCTT CARCACCTCA GCCACAGGGG AGGTGGAGGG	3120
GGATGTTGAG AGGGGAAGGA GGCAGAGCTG GGCAGCTGATCA CGTCAGACTC	3180
GGACAGGAAG GCAGGCAGAA GCTGGGTGCT CCTTATACAA CCCTACAGCC ACATTTTCCT	3240
CAGGACCGAG AGCCACAATG CTTCAGGAAA GCTAAAA CAATGCAGAG ACTAACTGGT	3300
TCCCTAAGCA TAGACAATGG CATTIGCCAA CACCAA CTCGAAAATG CCAGAGATTG	3360
GGTAGCTTTT GCCTGGCATT CAAAAACIGG GCTGACTGACTGCT GACAGCACGG	3420
TTAAACTTTT CACCCTGACC AGCACCCCAC AGACACCCCA ATAGTCTATG AGCAAGAAAG	3480
AGTGACCTGC AGCGCAGGGG AGGAGAAGAA AGCAGGGAM MAMAGTCCAC GTGATCCTGG	3540
ACAGATTCAT TCAAGGGCAG TGGGAATIGA CAAAAAMCA GTTCAAGGGA AGTCGGGAGA	3600
GTTCTAGGAG GCAGGGCTAT ATTGIGGGGG GAAGTCACAAG	3660
CCTGATTTCT AATACTATAT TITTCCTTA COMMONTANT TAGGAACTCT TTTTCTCTGT	3720
GTAGTAACTG AGGCTGTAAG ATTACTTAGT COMMENT ACAG GAAGAAACA TTCCTAAGAG	3780
GGAGTTAGCA GCACAAGGGC AATCCCCTTT COMCACMATA TGATTGGTTT TTTGAAAAAT	3840
TAAAGCCAAA CAGATTCAAG CCTAGGTC11 GCTAGCCCTT TGGTCAGCTG	3900
CATTTCAGCG ATGTTTACTA TCTGATTCAG COMMON CCCMT ALCTGCAGAA CCAATCAAAT	3960
TAAACAAACA CCCATTTGTA AATGTCTCAR MCMCCACTCT CGAGGTGAGT CTGCCAGGGC	4020
AAGAATAGAA TCTTTAGAGC AAACIGITTA ACACTTATTAAC AACATAAAGT	4080
AGTTTGGAAA TATTTACTTC ACAAGTATIG MENAACTTAC TTCTGACAGT TTTGGTATAT	4140
TGCTCAAAGG CAATCATTAT TTCAAGTGGC TTAAAGCAG	4200
TTATTGGCTA TTGCCATTTG CTTTTTGTT TOLEROWS A TGAGGAAAA ATTACGTTTT	4260
GGATTATTAA CCTACAGTCC AGAAAGCCTG IGAATTATTACC ATTGCCAATA GAGCCATAAA	4320
TATTTTTACC ACCTTCTAAC TAAATITAAC TOOTCAMUAC CAATAGAAAT CACAGACATT	4380
CTCAAAGTGG TAATAAGAGT ACCIGIGATI TICTLACTCA AATTTTTATA CTCAAAACTA	4440
TTATACTATA TTACAGTTGT TGCAGGTACG TTATATAAAAA CATGTTTAAA	4500
CTTTGAAATT AGACCTCCTG CTGGATCTTC TO TO TO THE TOTAL TCTATATTTT	4560
ATTTTGATAT TTTGATAATC ATATTICALL TTCGCCACT TTCACCAATG AGGTTCTTGG	4620
ATATATTGA AAACATCTTT CTGAGAAGAG ICCCCTAAC ATTGACATTG GTGCCTGAGA	4680
CATGCACACA CACAGAGTAA GAACTCCCAA AGAGCCIAAC TTGTTTTAAA GCTAGGGGTG TGCAAGACTG AAATTAGAAA GTTCTCCCAA AGATACACAG TTGTTTTAAA GCTAGGGGTG TGCAAGACTG AAATTAGAAA GTTCTCCCAG GAGCCTGGTA GGGTGCTGTC	4740
TGCAAGACTG AAATTAGAAA GTTCTCCCAA COMCMCCCTC GAGCCTGGTA GGGTGCTGTC	4800
AGGGGGGAAA TCTGCCGCTT CTATAGGAAT GCTCCTCTAAA GGACTTGTTT	4860
CTTGTGTTCT GGCTGGGTGCACG TTCTCAATGA GTTTGCAGAG	4920
GGATCTCCAG TTCCTAGCAT AGTGCCTGGC ACAGGCACAC CAGCACACCA GTAGTCCTGG	4980
TGAATGGAAA TATAAACTAG AAATATATA TOTTOTOTOTOTOTOTAAA ACCAGGTGGA	5040
TGTAAGTGTG TGTACGTGTG TGTGTGTGTGTGTGTGTGTGTTCTTT TTAAAAAGAA	5100
GATATAGGAA CTATTATTGG GGTATGGGTG CATAGATGT TCCTCCCAGC GTGAAGGCAA	5160
ACTCCAAACA GACTTCTGGA AGGTTATTTT COMOCGCACG MCTGTCTTCC CCCATGAAGG	5220
ACTCCAAACA GACTTCTGGA AGGTTATTTT CTAAGARICT IGCIGGCTCC CCCATGAAGG CCCCCCTGTG CACAGCCCCA CCCAGCCTCA CGAGCTCAC CATGAGCCAC CAAGGCCACC	5280
COMCOCACO CAGTATATAT AAACCICICI GGAGGIGGG	5304
CATCCAGGCA CCTCTCAGCA CAGC	

(2) INFORMATION FOR SEQ ID NO:3:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 6169 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

			GTATGTCTTT	GGCAGGATGA	TAAAGAATCA	120
TCCTATAAAC	TGTATAGCCT	CCATTCGGAT	TCCAGGCTGT	GTCTGCTCTT	ATTTTAGTGA	180
GGAAGAAGGA	GTATCCACGT	TAGCCAAGTG	CACCAAACAT	CACATCCAAT	ATGGTAAATC	240
CAGATGTTGC	TCCTGACAGA	AGCTATICTI	GATGGGCACT	TGCCCAAGGA	AAAATGCCAG	300
CATCAAACAG	GAGCTAAGAA	ACAGGAATGA	TCCCTTTGTT	TTTAATTTCA	GGAAAAAATG	360
GAGAGCAAAT	AATGATGAAA	AMTMARCIII	CAGCTCAGAA	AAAAGATGTT	TCCAAATTGG	420
ATGAGGACCA	AAATCAATGA	CCCAACACAC	CTCCATGTGA	GCTTGATGGG	AAAATGGGAA	480
TAATTAAGTA	TITGTTCCTT	CAMCAGAGAC	CAAAGTGGAT	TATTATTTA	AAAACCAGAT	540
AAACGTCAAA	AGCATGATCT	ACMMCAGAICC	GGTCATGTTA	GCAAAGGACA	TAACAATAAC	600
GGCATCACTC	TGGGGAGGCA	AGTICAGGAA	GAAAATGGGG	ACTGGGAAAG	CTTTCATAAC	660
AGCAAAATCA	AAATTCCGCA	MATGCAGGAG	ACCTCCCCGT	CTATACCAGG	GAACACAAAA	720
AGTGATTAGG	CAGTTGACCA	A COUNTRY A CO	GAAATATGAA	AAACTGAGAG	CAAAACAAAA	780
ATTGACTGGG	CTAAGCCIGG	ACTITICARGO	GAGCCTTCAA	AGCAGCAGTG	CCCCTCAGCA	840
GACATGGTTA	AAAGGCAACC	MGAACAIIGI	CCAGTTTTCT	TAAGGAATCT	TAAGAAACTC	900
GGGACCCTGA	GGCATTTGCC	ACCAMMMAQ	GTATAAAACA	AATATGCGAT	GCATAATCAG	960
TTGAAAGATC	ATGAATTITA	MUCKILLIA:	AGGCATACAA BETCCCCATC	GGATAACGTG	TCCCAGCTCC	1020
TTTAGACATG	GGTCCCAATI	CANADORGE	TGTCCCCATC	CTAACTTTTT	CAGAATGATC	1080
GGATAGGTCA	GAAATCATTA	CCCCCGATGT	GTCTGACCTA	CAACCACATC	TACAACCCAA	1140
TGTCATAGCC	CTCACACACA	CTCTCATCT	AGTAGGTCCC	ATTACAAATC	CCACCTCCCC	1200
GTGCCTCAAC	CATTGTTAAC	ACAGGAAGT	TCCCCACTCT TCCCCACTCTA	AGACTTCTGC	ATCACGATGT	1260
TGTGCAGCCC	ATCCCGCTCC	CCCTCAGGG	TCCCCACTC1 TCTGTGTCTTA	CACCTACCTC	TATGCTCTAC	1320
TACAGCCAGA	AGCTCCGIGA	CTGCCTCCC	A GGTTCAAGCA	ATTCTCCTGT	CTCAGCCTCC	1380
ACCTGAGCT	ACTGUARCE	CECACECCC	GCTAATTTT	GTATTGTTAC	TAGAGATGGG CACCCACCTC	1440
CGCGTAGCT	GGACTACAG	, CGCACGCCC.	A ACTCCTGACC	TCAGGTGAT	CACCCACCTC A GGGTCAGTGT	1500
GTTTCACCA'	P ATTAGECEG	TTACAGGCA	T GAGTCACCGC	GCCCGGCCA	A GGGTCAGTGT A AAAGCTGTGA	1560
AGCCTCCTA	A AGIGCIGGG	TGGTTTACT	A AACCAACAGG	GAAACAGAC	A AAAGCTGTGA T AGTCCCAGAC	1620
TTAATAAGG	A ATAACIIGA	2 ATGGGGAAT	G GTGCCATGAC	CTGCCTGCC'	r AGTCCCAGAC A CCATTTTATT	1680
TAATTTCAG	G GATTCIIGG	C TTCCCTCAT	C CTCATTTTC	GGCTAAGTT	A CCATTTTATT A CATAAACTAG	1740 1800
CACTGGTCC	T CHICACIII	G CCTCCACAT	C GTTACTGAAA	TAAGAGTAT	A CATAAACTAG C AGAGACTCCT	1860
CACCATGCT	C CCCCATCTG	T GTGTGTGTA	T AGGGGAGGA	GGCATACCC	C AGAGACTCCT A CCCGGGGTCC	1920
TOLLOGO	C CCCAGAGGT	T TCCTCTCCA	G CTGGGGGAG	CCTGCAAGC	A CCCGGGGTCC C ACTCTCTAGG	1980
TGAAGCCCC	T GAGCAACCT	G CCAGCCCGT	G CCACTGGTT	3 TTTTGTTAT	C ACTCTCTAGG C ATTGACAATT	2040
TGGGTGTCC	C TOTAL TAREST	C TGTGTGACT	C GTTCATTCA	r ccaggcatt	C ATTGACAATT G CAGTCACTGC	2100
GACCIGIIG	C TTATATCTG	C CAGACACCA	G AGACAAAAT	G GTGAGCAAA	G CAGTCACTGC A TTAATAGCCA	2160
TATIGAGIA	C TEGAGGTGA	C AGTTTCTCA	T GGAAGACGT	G CAGAAGAAA	A TTAATAGCCA G AATTGTGCGC	2220
CCTACCTTC	A ACCCAGTGO	T GAAAGAAAC	G AAATAAACA	C CATCTTGAA	G AATTGTGCGC G TGCCCGGAGG	2280
ACCARCT III	T AACAAGGCC	A CCTCCCTAC	C GCCCCCTGC	T GCCTCCATC	G TGCCCGGAGG G CAGCTGGCCT	2340
CCCCCAAGC	C CGAGTCTTC	C AAGCCTCC	C CTCCATCAG	T CACAGOGCI	G CAGCTGGCCT	2400
GCCTCGCTT	C CCGTGAATO	G TCCTGGTG	A TOTGAGOTG	G AGACTCCTT	G GCTCCAGGCT G ACAGTGTTTC	2460
CCAGAAAGO	A AATGGAGAG	G GAAACTAG	IC TAACGGAGA	A TOTGGAGGG	G ACAGTGTTTC A CTGCAGGGAG	2520
CTCAGAGG	A AAGGGGCC	C CACGTCCA	GG AGAATTCCA	G GAGGTGGG	A CTGCAGGGAG G CAAGGCTGAA	2580
TGGGGACG	T GGGGCTGA	SC GGGTGCTG	AA AGGCAGGAA	G GTGAAAAGG	G CAAGGCTGAA T TCCTGTGAGC	2640
GCTGCCCA	GA TGTTCAGT	T TGTTCACG	GG GCTGGGAG1	TICCGIIG	T TCCTGTGAGC	2700
	~~ ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	TO TIGGAGGAG	AA GAAGTCTAT	T TCATGAAG	GG ATGCAGTTTC GA AGGCCTTTAT	2760
ATAAAGTC	AG CTGTTAAA	AT TOCAGGGT	GI GCAIGGGIA	መ መአአመጥሮአሮ(C AAGAAGTGAC	2820
TTAATGGG	AA TATAGGAA	GC GAGCTCAT	TI CCIAGGCCC	C CCTCTGGT	G ACTTGGCTTA	2880
TGGAGTCT	TT TCTTTCAT	GT CTTCTGGG	CA ACIACICAC	20 CCTCTCC	GG TTCTGCCATT GT CTTCTCTGCT	2940
TGCAAGAC	GG TCGAAAAC	CT TGGAATCA	om concommece	TO COCCOATA	CTTCTCTGCT	3000
GGTTGGCT	GT GCGACCGT	GG GCAAGIGI	CT CICCITOO	C CTCTCATT	CT CTGTGAGGGG	3060
ATAAAGAC	CC TTGCAGCT	CT CGTGTTC1	GI GRACACITA	CA GCCACAGG	GG AGGTGGAGGG	3120
GGATGTTG	AG AGGGGAAG	GA GGCAGAGC	TO CAMOACH	CC TCACTGAT	CA CGTCAGACTC	3180
GGACAGGA	AG GCAGGCAG	AA GCTGGGTG	CI CCAICAGI	AA CCCAACAG	CC ACATTTTCCT	3240
CAGGACCG	AG AGCCACAA	TG CTTCAGGA	TAN GCICUMIO	AN CANTICAG	AG ACTAACTGGT	3300
TCCCTAAG	CA TAGACAAT	GG CATTIGCO	MA IMACCAM	AA GTGGAAAA	TG CCAGAGATTG	3360
GGTAGCTT	TT GCCTGGCA	TT CAAAAAC	TO COLCADO	CC ACTGACTG	CT GACAGCACGG	3420
TTAAACTT	TT CACCCTGA	ICC AGCACCC	AC GCAGGIOA	CC ATACTOTA	TG AGCAAGAAAG	3480
AGTGACCT	IGC AGCGCAGG	GG AGGAGAA	JAA AAAGAGAGAG	አመ መእጥልርጥርር	AC GTGATCCTGG	3540
ACAGATTO	AT TCAAGGG	AG TGGGAAT	TGA CCACAGGO	CA CUMCAAGO	CA AGTCGGGAGA	3600
GTTCTAGG	BAG GCAGGGC	PAT ATTGTGG	MAN CANCOUCY	CT AATTCTC	GC AAGTCACAAG	3660
CCTGATT	rct aatacta:	PAT TITTCCT	IIM CAMBCION	AM MAGGAAC	CT TTTTCTCTGT	3720
GTAGTAA	CTG AGGCTGT	AAG ATTACTT	WGI IICICCII	NC CAACAAA	CA TTCCTAAGAG	3780
GGAGTTA	GCA GCACAAG	GGC AATCCCG	TIL CITITUM	שא שכאששכני	PTT TTGAAAAAT	3840
TAAAGCC.	AAA CAGATTC	AAG CCTAGGT	CII GCIGACIA	AC TAGTACC	TTT TGGTCAGCTG	3900
CATTTCA	GCG ATGTTTA	CTA TCTGATT	CAG AAAAIGA	TO A A CTGCA	CAATCAAAA	3960
TAAACAA	ACA CCCAGTT	GTA AATGTCT	CAA GIICAGO	nom concerc	AGT CTGCCAGGGC	4020
AGAATAG	AAT CTTTAGA	GCA AACTGTG	TTO ACACTOR	TGT TGGTATT	AAC AACATAAAGT	4080
AGTTTGG	AAA TATTTAC	TIC ACAAGTA	TITO MCMC+O+			

		TTCAAGTGGC	TTAAAGTTAC	TTCTGACAGT	TTTGGTATAT	4140
1001010	CLUST COLD	CTTTTTGTTT	TTTCTCTTTG		TGTAAAGCAG	4200
	TTGCCATTTG	AGAAAGCCTG	TGAATTTGAA		ATTACATTTT	4260
00112 2272 27	CCTACAGTCC	TAAATTTAAC	ATTTTATTCC		GAGCCATAAA	4320
TGTTTTTACC	ACCTTCTAAC	ACCTGTGATT	TTGTCATTAC		CACAGACATT	4380
CTCAAAGTGG	TAATAACAGT	TGCAGATACG	TTGTAAGTGA	AATATTTATA	CTCAAAACTA	4440
TTATACTATA	TTACAGTTGT		TTTTTAACAT	ATTAATAAAA	CATGTTTAAA	4500
CTTTGAAATT	AGACCTCCTG	CTGGATCTTG ATATTTCATT	ATCATTTGTT	TCCTTTGTAA	TCTATATTTT	4560
ATTTTGATAT	TTTGATAATC		TTCCCCAGAT	TTCACCAATG	AGGTTCTTGG	4620
ATATATTTGA	AAACATCTTT	CTGAGAAGAG	AGAGGCTAAC	ATTGACATTG	GTGCCTGAGA	4680
CATGCACACA	CACAGAGTAA	GAACTGATTT	AGATACACAG	TTGTTTTAAA	GCTAGGGGTG	4740
TGCAAGACTG	AAATTAGAAA	GTTCTCCCAA	GCTCTCCCTG	GAGCCTGGTA		4800
AGGGGGGAAA	TCTGCCGCTT	CTATAGGAAT	TGTCCCTGCT	ACGTCTTAAA	GGACTTGTTT	4860
CTTGTGTTCT	GGCTGGCTGT	TATTTTTCTC	ACAGTGCAGG	TTCTC A ATGA	GTTTGCAGAG	4920
GGATCTCCAG	TTCCTAGCAT	AGTGCCTGGC	TTGTTGAAAT	CAGCACACCA		4980
TGAATGGAAA	TATAAACTAG	AAATATATCC		TGTAAAACCA		5040
TGTAAGTGTG	TGTACGTGTG	TGTGTGTGTG		TTCTTTTTAA		5100
TAGGAACTAT	TATTGGGGTA	TGGGTGCATA				5160
CAAACAGACT	TCTGGAAGGT					5220
CCTGTGCACA	GCCCCACCCA					5280
GCTCCCCAGT	ATATATAAAC		· · · · · · · · · · · · · · · · · · ·		TCTGCAATGA	5340
CAGGCACCTC	TCAGCACAGC				CAGCTGCTGC	5400
GGTTCTTCTG						5460
TTCTGGCCTC	CCTGGTGTGC					5520
ACCAGAGTGG						5580
CAGAGCAGAG						5640
GCTTAGACCT	GGAGGCCACC					5700
TGACCTTGG						5760
GCACCCTGAG						5820
ACAGCAACC'	r cctccgaga					5880
AAAATGAGA	A TCTGGCCAG					5940
GGGGCCAGT	G TCCCCAGAC					6000
GTAAGAATG	C AGAGTGGGG					6060
CTACAGGCG						6120
AGATGAATT						6169
AGCTTCATT	T AGATTAGTG	G TTCAGAGTT	C TIGIGCECE			

(2) INFORMATION FOR SEQ ID NO:4:

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 926 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single

- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

AAGGTAGGCA CATTGCCCTG CAATTTATAA TTTATGAGGT GTTCAATTAT GGAATTGTCA AAAGTAGAGA GACTACAATG TAGCCATAAC TCAGGCCCAA TCATGTTTTA TCTTTCTT TCTTTCTTT TCTTTCTTT TCTTCTTCTT	60 120 180 240 300 360 420 480 540 660 720 780 900 926
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(2) INFORMATION FOR SEQ ID NO:5:

(i) SEQUENCE CHARACTERISTICS:

PCT/US00/00559 WO 00/42220

- (A) LENGTH: 2099 base pairs (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

			mamamamma.	GAAGATTATG	GATTAAGTGG	60
TGGCTCTGCC	AAGCTTCCGC				AGGAGAGCCT	120
TGCTTCGTTT	701111	7 7 7110 0110 011			GCGAGACCCC	180
CTCACGCTGA	CARACTIC CO.	Wichman			TGGCACGGAT	240
AAGCCCACCT	ACCCCTACAC		VCO 1 CO. ICI.		CCCTTCTAAG	300
GTCCGCCAGG	TTTTTGAGTA	20	AGCCAGTTTA		GGGGAGCCTC	360
GTTCACATAC	TGCCTAGGCC	ACTGGAAAGC	ACGGGTGCTG	AGCTGAATAC	CGAGACAGTG	420
TATTTCCAGG	GCGCTGAGTC	CAGAACTGTC	ATAAGATATG	AGTTCCCGTA	TTCTTGGGGT	480
AAGGCTGAGA	AGGAAATCCC	TGGAGCTGGC	TACCACGGAC	TCTGGGTCAT	TTACAGCACC	540
GGCTACACGG	ACATTGACTT	GGCTGTGGAT	GAAGCAGGCC	CAGAGAATCT	GGAACTCGAA	600
GATGAGGCCA	AAGGTGCCAT	TGTCCTCTCC	AAACTGAACC	ATGCCTTCAT	CATCTGTGGC	660
CAAACCTGGG	AGACAAACAT	CCGTAAGCAG	TCAGTCGCCA GCAGATGCTA	CCGTCAACTT	TGCTTATGAC	720
ACCTTGTACA	CCGTCAGCAG	CTACACCTCA	ATCCCATTCA	AGAACCGCTA	TAAGTACAGC	780
ACAGGCACAG	GTATCAGCAA	GACCCTGACC		CCTGGGACAA	CTTGAACATG	840
AGCATGATTG		CCTGGAGAAG	AAGCTCTTTG TGAAAAGCCT	CCAAGCTGTA	CAGGCAATGG	900
GTCACTTATG		CTCCAAGATG	AGCAGGCTGA		AGCCAGCCAG	960
CAGAAGGAGA		TCCTGGGGGG	TTTCATTAAT		GAACATGGTC	1020
GGCCCAGGCA	GCTTTGACTG	CTTTCCAAGT	AGGGCGTAGA		TAATAAATAT	1080
ACCATCTAAC			TTTAATGACA		TTTCTTGTGA	1140
CCTTTATCTT	CTGTCAGCAT				TCTTGCATGT	1200
TTTGGGGCA					GAATAGCTCC	1260
TACATGGTTA						1320
TCTGGCCAG						1380
ATAGAATAC						1440
CGTTCTCTTC						1500
ATCAGGTAA!						1560
GTGTGTGTG'						1620
CGCCAAGCA						1680
CCGGAGCAA'			·			1740
AGTTTGCTT			·		CTCCCAGGTT	1800
CACCCAGGC					TGCCACCACG	1860
CACACCATT					CAGGATGGTC	1920
CCTGGCTAA		•			GATTACAGGC	1980
TCGATCTCC					GGTATGAATT	2040
GTGAGCCAC			·			2099
TTATAAGCC	A CACCTCAGG	I GGWGWWWGC	1 10111001111	- · · ·		

- (2) INFORMATION FOR SEQ ID NO:6:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 19 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

TGAGGCTTCC TCTGGAAAC

(2) INFORMATION FOR SEQ ID NO:7:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 base pairs
 (B) TYPE: nucleic acid

 - (C) STRANDEDNESS: single (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
TGAAATCAGC ACACCAGTAG	20
(2) INFORMATION FOR SEQ ID NO:8:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	21
GCACCCATAC CCCAATAATA G	21
(2) INFORMATION FOR SEQ ID NO:9:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	2.2
AGAGTTCCCC AGATTTCACC	20
(2) INFORMATION FOR SEQ ID NO:10:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:	20
ATCTGGGGAA CTCTTCTCAG	20
(2) INFORMATION FOR SEQ ID NO:11:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	1.0
TACAGTTGTT GCAGATACG	19
(2) INFORMATION FOR SEQ ID NO:12:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
() SEQUENCE DESCRIPTION: SEQ ID NO:12:	

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TO THE TOTAL OF THE CONTRACT O	21
ACAACGTATC TGCAACAACT G	
(2) INFORMATION FOR SEQ ID NO:13:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	20
TCAGGCTTAA CTGCAGAACC	20
(2) INFORMATION FOR SEQ ID NO:14:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	. 19
TTGGTTCTGC AGTTAAGCC	13
(2) INFORMATION FOR SEQ ID NO:15:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	19
AGCAGCACAA GGGCAATCC	19
(2) INFORMATION FOR SEQ ID NO:16:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:	18
ACAGGGCTAT ATTGTGGG	
(2) INFORMATION FOR SEQ ID NO:17:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:	
CCTCA CATCC CACCTGTCC	19

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18: CTGAAGCATT AGAAGCCAAC (2) INFORMATION FOR SEQ ID NO:19: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19: ACCTTGGACC AGGCTGCCAG (2) INFORMATION FOR SEQ ID NO:20: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20: AGGTTTGTC GAGTTCCAG (2) INFORMATION FOR SEQ ID NO:21: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	(A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18: CTGAAGCATT AGAAGCCAAC (2) INFORMATION FOR SEQ ID NO:19: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19: ACCTTGGACC AGGCTGCCAG (2) INFORMATION FOR SEQ ID NO:20: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20: AGGTTTGTTC GAGTTCCAG (2) INFORMATION FOR SEQ ID NO:21: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21: ACAATTACTG GCAAGTATGG (2) INFORMATION FOR SEQ ID NO:22: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	(2) INFORMATION FOR SEQ ID NO:18:	
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(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21: ACAATTACTG GCAAGTATGG	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21: ACAATTACTG GCAAGTATGG (2) INFORMATION FOR SEQ ID NO:22: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	AGGTTTGTTC GAGTTCCAG	17
(A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21: ACAATTACTG GCAAGTATGG	(A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21: ACAATTACTG GCAAGTATGG (2) INFORMATION FOR SEQ ID NO:22: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	(2) INFORMATION FOR SEQ ID NO:21:	
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	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:		
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	19	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	
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(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:	
ACACCTCAGC AGATGCTACC	20
(2) INFORMATION FOR SEQ ID NO:24:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
ATGGATGACT GACATGGCC	19
(2) INFORMATION FOR SEQ ID NO:25:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	19
AAGGATGAAC ATGGTCACC	
(2) INFORMATION FOR SEQ ID NO:26:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1548 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: CDNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	
AGAGCTTTCC AGAGGAAGCC TCACCAAGCC TCTGCAATGA TGCAGCTTTG GGCCTGAGAT GGCCTGAGAT GATGTGGGGG CCAGGACAGC TCAGCTCAGG TCAGCTCAGG TCAGCTCAGG TCAGCTCAGG TCAGCTCAGG TCAGCTCAGG TCAGCTCAGG TCAGCTCACC AAACCTTACA AAACCTCACC AAAGCTCAGC AAACCCAAC GGAGACACC AAACCCAAC GGAGACCCCAGG AGACCCAGGA GACCCAGGA GACCCAGGA AAACCCAAAC AAACCCAAC CCTACAGAAC AAAACTAACA ACCCCAGC TCCAGAGAAC TCCAGAGAAC TCCCCAGAC TCCAGAGAC TCCCCAGAC TCCAGAGAC TCCCCAGAC TCCAGAGAC TTCTCACGAC TCCCAGACC TCCAGACAC TCCAGACC TCCAGACAC TCCAGACAC TCCCAGACC TCCAGACAC TCCAGACAC TCCCAGACC TCCAGACAC TCCAGAGAC TCCCAGACC TCCAGACAC TCCCAGAC TCCCCAGC TCCCAGAC TCCCCAGC TCCCAGAC TCCCCAGC TCCCAGAC TCCCAGAC TCCCCAGC TCCCAGAC TCCCCAGC TCCCAGAC TCCCCAGC TCCCAGAC TCCCAGAC TCCCCAGC TCCCAGAC TCCCCAGC TCCCAGAC TCCCCAGC TCCCAGAC TCCCAGC TCCCAGAC TCCCCAGC TCCCAGAC TCCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCAGAC TCCCCAGC TCCCAGC TCCCAGC TCCCAGAC TCCCCAGC TCCCAGAC TCCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCACC TCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCAGC TCCCACC TCCCAGC TCCCCAGC TCCCAGC TCCCCAGC TCCCCAGC TCCCCAGC TCCCCAGC TCCCCAGC TCCCCAGC TCCCCAGC TC	60 120 180 240 300 420 420 660 720 780 960 1020

TTCCCGTATT TGGGTCATTT GAGAATCTGG GCCTTCATCA GTCAACTTTG	ACAGCACCGA AACTCGAACA TCTGTGGCAC	TGAGGCCAAA AACCTGGGAG CTTGTACACC AGGCACAGGT	GGTGCCATTG ACAAACATCC GTCAGCAGCT ATCAGCAAGA TACAACCCCC	TCCTCTCAA GTAAGCAGTC ACACCTCAGC CCCTGACCAT TGGAGAAGAA	CCACGGACAG AGCAGGCCTC ACTGAACCCA AGTCGCCAAT AGATGCTACC CCCATTCAAG GCTCTTTGCC	1140 1200 1260 1320 1380 1440 1500
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(2) INFORMATION FOR SEQ ID NO:27:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 178 amino acids
 (B) TYPE: amino acid

 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

Thr Gly Ala Val Val Tyr Ser Gly Ser Leu Tyr Phe Gln Gly Ala Glu 10 Ser Arg Thr Val Ile Arg Tyr Glu Leu Asn Thr Glu Thr Val Lys Ala 25 Glu Lys Glu Ile Pro Gly Ala Gly Tyr His Gly Gln Phe Pro Tyr Ser 40 Trp Gly Gly Tyr Thr Asp Ile Asp Leu Ala Val Asp Glu Ala Gly Leu 55 Trp Val Ile Tyr Ser Thr Asp Glu Ala Lys Gly Ala Ile Val Leu Ser 70 75 Lys Leu Asn Pro Glu Asn Leu Glu Leu Glu Gln Thr Trp Glu Thr Asn Ile Arg Lys Gln Ser Val Ala Asn Ala Phe Ile Ile Cys Gly Thr Leu 100 105 110 85 Tyr Thr Val Ser Ser Tyr Thr Ser Ala Asp Ala Thr Val Asn Phe Ala Tyr Asp Thr Gly Thr Gly Ile Ser Lys Thr Leu Thr Ile Pro Phe Lys 135 Asn Arg Tyr Lys Tyr Ser Ser Met Ile Asp Tyr Asn Pro Leu Glu Lys Lys Leu Phe Ala Trp Asp Asn Leu Asn Met Val Thr Tyr Asp Ile Lys 150 170 165 Leu Ser

(2) INFORMATION FOR SEQ ID NO:28:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 131 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

Arg Phe Asp Leu Lys Thr Glu Thr Ile Leu Lys Thr Arg Ser Leu Asp Tyr Ala Gly Tyr Asn Asn Met Tyr His Tyr Ala Trp Gly Gly His Ser 25 Asp Ile Asp Leu Met Val Asp Glu Ser Gly Leu Trp Ala Val Tyr Ala 40 Thr Asn Gln Asn Ala Gly Asn Ile Val Val Ser Arg Leu Asp Pro Val 55 Ser Leu Gln Thr Leu Gln Thr Trp Asn Thr Ser Tyr Pro Lys Arg Xaa

(2) INFORMATION FOR SEQ ID NO:29:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 178 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:

Gly Thr Gly Gln Val Val Tyr Asn Gly Ser Ile Tyr Phe Asn Lys Phe 10 Gln Ser His Ile Ile Ile Arg Phe Asp Leu Lys Thr Glu Thr Ile Leu 25 Lys Thr Arg Ser Leu Asp Tyr Ala Gly Tyr Asn Asn Met Tyr His Tyr 45 40 35 Ala Trp Gly Gly His Ser Asp Ile Asp Leu Met Val Asp Glu Asn Gly 55 60 Leu Trp Ala Val Tyr Ala Thr Asn Gln Asn Ala Gly Asn Ile Val Ile 65 70 75 80 Ser Lys Leu Asp Pro Val Ser Leu Gln Ile Leu Gln Thr Trp Asn Thr 90 Ser Tyr Pro Lys Arg Ser Ala Gly Glu Ala Phe Ile Ile Cys Gly Thr 85 105 100 Leu Tyr Val Thr Asn Gly Tyr Ser Gly Gly Thr Lys Val His Tyr Ala 125 120 Tyr Gln Thr Asn Ala Ser Thr Tyr Glu Tyr Ile Asp Ile Pro Phe Gln 140 135 Asn Lys Tyr Ser His Ile Ser Met Leu Asp Tyr Asn Pro Lys Asp Arg 130 155 150 Ala Leu Tyr Ala Trp Asn Asn Gly His Gln Thr Leu Tyr Asn Val Thr 170 165 Leu Phe

(2) INFORMATION FOR SEQ ID NO:30:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 177 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

Trp Val Ile Phe Thr Thr Glu Lys Ser Ala Gly Lys Ile Val Val Gly Lys Val Asn Val Ala Thr Phe Thr Val Asp Asn Ile Trp Ile Thr Thr 70 85 Gln Asn Lys Ser Asp Ala Ser Asn Ala Phe Met Ile Cys Gly Val Leu 105 Tyr Val Thr Arg Ser Leu Gly Pro Lys Met Glu Glu Val Phe Tyr Met 100 120 Phe Asp Thr Lys Thr Gly Lys Glu Gly His Leu Ser Ile Met Met Glu 140 135 Lys Met Ala Glu Lys Val His Ser Leu Ser Tyr Asn Ser Asn Asp Arg 155 150 Lys Leu Tyr Met Phe Ser Glu Gly Tyr Leu Leu His Tyr Asp Ile Ala 170 165 Leu

(2) INFORMATION FOR SEQ ID NO:31:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 74 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: None
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

Gly Val Val Tyr Ser Arg Leu Thr Glu Thr Leu Ala Gly Tyr Asn Asn 10 15

Tyr Ala Trp Gly Gly Asp Ile Asp Leu Val Asp Glu Gly Leu Trp Tyr 25 30

Thr Ala Gly Ile Val Ser Lys Leu Pro Leu Gln Thr Trp Thr Lys Ala 35

Phe Ile Ile Cys Gly Thr Leu Tyr Val Thr Tyr Val Tyr Ala Tyr Thr 50

Ile Tyr Asp Tyr Asn Pro Lys Leu Tyr Leu 65

- (2) INFORMATION FOR SEQ ID NO:32:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 504 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (V) FRAGMENT TYPE: N-terminal
- · (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

 Met Arg Phe Phe Cys Ala Arg Cys Cys I
 Ser Phe Gly Pro Glu Met Pro 10 15

 Ala Val Gln Leu Leu Leu Leu Leu Ala Cys Leu Val Trp Asp Val Gly Ala 20 20 30

 Arg Thr Ala Gln Leu Arg Lys Ala Asn Asp Gln Ser Gly Arg Cys Gln 30 40

 Tyr Thr Phe Ser Val Ala Ser Pro Asn Glu Ser Ser Cys Pro Glu Gln 55 60

 Ser Gln Ala Met Ser Val Ile His Asn Leu Gln Arg Asp Ser Ser Thr 70 70 70 75 80

 Gln Arg Leu Asp Leu Glu Ala Thr Lys Ala Arg Leu Ser Ser Leu Glu 90 95

 Ser Leu Leu His Gln Leu Thr Leu Asp Gln Ala Ala Arg Pro Gln Glu Glu 100

```
Thr Gln Glu Gly Leu Gln Arg Glu Leu Gly Thr Leu Arg Arg Glu Arg
                             120
Asp Gln Leu Glu Thr Gln Thr Arg Glu Leu Glu Thr Ala Tyr Ser Asn
      115
                         135
                                             140
    130
Leu Leu Arg Asp Lys Ser Val Leu Glu Glu Glu Lys Lys Arg Leu Arg
                                         155
                    150
Gln Glu Asn Glu Asn Leu Ala Arg Arg Leu Glu Ser Ser Ser Gln Glu
                                     170
                165
Val Ala Arg Leu Arg Arg Gly Gln Cys Pro Gln Thr Arg Asp Thr Ala
180 185 190
Arg Ala Val Pro Pro Gly Ser Arg Glu Val Ser Thr Trp Asn Leu Asp
     195
Thr Leu Ala Phe Gln Glu Leu Lys Ser Glu Leu Thr Glu Val Pro Ala
                                              220
                        215
Ser Arg Ile Leu Lys Glu Ser Pro Ser Gly Tyr Leu Arg Ser Gly Glu 225 235
    210
Gly Asp Thr Gly Cys Gly Glu Leu Val Trp Val Gly Glu Pro Leu Thr
                 245
Leu Arg Thr Ala Glu Thr Ile Thr Gly Lys Tyr Gly Val Trp Met Arg 260 265 270
 Asp Pro Lys Pro Thr Tyr Pro Tyr Thr Gln Glu Thr Thr Trp Arg Ile
275 280 285
 Asp Thr Val Gly Thr Asp Val Arg Gln Val Phe Glu Tyr Asp Leu Ile
                                              300
                          295
 Ser Gln Phe Met Gln Gly Tyr Pro Ser Lys Val His Ile Leu Pro Arg
320
     290
 Pro Leu Glu Ser Thr Gly Ala Val Val Tyr Ser Gly Ser Leu Tyr Phe 325
 Gin Gly Ala Glu Ser Arg Thr Val Ile Arg Tyr Glu Leu Asn Thr Glu
                                 345
            340
 Thr Val Lys Ala Glu Lys Glu Ile Pro Gly Ala Gly Tyr His Gly Gln
 Phe Pro Tyr Ser Trp Gly Gly Tyr Thr Asp Ile Asp Leu Ala Val Asp 370 380
 Glu Ala Gly Leu Trp Val Ile Tyr Ser Thr Asp Glu Ala Lys Gly Ala
385 390 395
 Ile Val Leu Ser Lys Leu Asn Pro Glu Asn Leu Glu Leu Glu Gln Thr
                                      410
                 405
 Trp Glu Thr Asn Ile Arg Lys Gln Ser Val Ala Asn Ala Phe Ile Ile
              420
 Cys Gly Thr Leu Tyr Thr Val Ser Ser Tyr Thr Ser Ala Asp Ala Thr 435
  Val Asn Phe Ala Tyr Asp Thr Gly Thr Gly Ile Ser Lys Thr Leu Thr
                                             460
                           455
      450
  Ile Pro Phe Lys Asn Arg Tyr Lys Tyr Ser Ser Met Ile Asp Tyr Asn 480
  Pro Leu Glu Lys Lys Leu Phe Ala Trp Asp Asn Leu Asn Met Val Thr
                                       490
                  485
  Tyr Asp Ile Lys Leu Ser Lys Met
               500
```

(2) INFORMATION FOR SEQ ID NO:33:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

CAAACAGACT TCCGGAAGGT

(2) INFORMATION FOR SEQ ID NO:34:

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 5271 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:

ልጥርጥጥጥርጥጥር	AGTTTACCTC	AGGGCTATTA '	TGAAATGAAA '	TGAGATAACC	AATGTGAAAG	60
MOOM3 M3 3 3 C	mcmamacccm	CCAMMCGGAM (GTATGTCTT	GGCAGGATGA	IWWWGWWICW	120
CC3 3 C3 3 CC3	CMMMCCMCCM	TACCCAACTG	TCCAGGCTGT	GICIGCICII	WITTINGTON	180
CA CA MORMOCO	mccmcacacaca	ልርርጥልጥጥር ጥ ጥ ፣	CAGGAAACAT	CACATCCAAT	ATGGTAAATC	240
ORMORRAGEO	CACCMAACAA	ACACCA ATCA	GATGGGCACT	TGCCCAAGGA	AAAAIGCCAG	300
0303003330	አአጥሮአጥሮልልል	<u> የ</u> ምምጋል ልጥጥጥ	TCCCTTTGTT	TTTAATTTCA	GGAAAAATG	360
1001001001	እ አ አ መረግ እ ጥር ጋ	ΑΠΝΑΓΓΑΝΑΑ	CAGCTCAGAA	AAAAGATGTT	TCCAAATIGG	420
ma amma acma	WIND CAUCAUT	CCGA AGAGAC	CTCCATGTGA	GCTTGATGGG	AAAATGGGAA	480
*******	አ ረ ረ አ መር አ ጥርጥ	ረልጥሮልርልጥሮሮ	CAAAGTGGAT	TATTATTTA	AAAACCAGAI	540
CCC A MC A CMC	mccccaccca	ACTITICACGAA	GGTCATGTTA	GCAAAGGACA	TAACAATAAC	600
100111100	አ አ አ መጥሮሮሮሮ እ	AATGCAGGAG	GAAAATGGGG	ACTGGGAAAG	CTTTCATAAC	660
3 CMC 3 MM3 CC	これで中でこれでごれ	ጥርጥጥርርርልልር	ACCTCCCCGT	CTATACCAGG	GAACACAAAA	720
3 MMC 3 CMCCC	ርመን አርርርጥርር	ልርጥጥጥሮልልር ር	GAAATATGAA	AAACTGAGAG	CAMMACAMAM	780
	******	አርአ አር አጥጥርጥ	GAGCCTTCAA	AGCAGCAGTG	CCCCTCAGCA	840
COCK CCCCCX	CCCAMMTCCC	ጥጥጥልርርልልርር	CCAGTTTTCT	TAAGGAATCT	TAAGAAACIC	900
	지 때 (가지 때 (대대) 지 지	א מידיידיידית אריר א	GTATIAAAACA	AATATUUGAT	GCWIWVICVO	960
mmma	CCMCCC3 3 ጥጥ	ጥጥልጥልልልርጥሮ	AGGCATACAA	GGATAACGTG	TUCCAGUTUC	1020
CORMICOMOR	<u> </u>	CAAATCACTG	TGTCCCCATC	CTAACTTTTT	CHGWWIGWIC	1080
momos macco	CTCACACACA	GGCCCGATGT	GTCTGACCTA	CAACCACATC	TACAACCCAA	1140
OMOGOMON NO	ር አ ጥጥር ጥጥ አ እ ር	CHCHC AUCUC	AGTAGGTCCC	A'I'I'ACAAA'IG	CCACCICCC	1200
momooa coco	A THE CHECK THE CHECK	ACAGGAAGTC	TCCCCACTCT	AGACTTCTGC	ATCACGATGI	1260
MA CACCCACA	えごごかごごところ	GGGTGAGGGT	CTGTGTCTTA	CACCTACCTG	TATGUTUTAL	1320
3 CCBC 3 CCBC	አርጥርር እአርርጥ	CTGCCTCCCA	GGTTCAAGCA	ATTCTCCTGT	CICAGCCICC	1380
COCCETA COTO	CCACTACACC	CGCACGCCCG	CCTAATTTTT	GTATTGTTAG	TAGAGATGGG	1440
	* ************************************	CTGGTCTTGA	ACTCCTGACC	TCAGGIGATC	CACCCACCIC	1500
* CCCMCCM3 3	A CITIC CITIC CCA	ጥጥልሮልርርርልጥ	GAGTCACCGC	GCCCGGCCAA	GGGTCAGIGI	1560
mma ama acca	አመአአሮሞሞርልል	T	AACCAACAGG	GAAACAGACA	AMAGCIGIGA	1620
maxmmmcacc		ATCCCCCA ATC	GTGCCATGAG	CTGCCTGCCT	AGICCCAGAC	1680
	፣ CX ጥCXCጥጥጥር	ጥጥርርርጥርልጥር	CTCATTTTCA	GGCTAAGTTA	CCATTTALL	1740
CA COMPCOMP	1 - መጥርጥርርጥ እ እር	CCTCCACATC	GTTACTGAAA	TAAGAGTATA	CATAAACTAG	1800
	, CCCCATCTCT	СТСТСТСТАТ	AGGGGAGGAG	GGCATACCCC	AGAGACICCI	1860
man naccent	*	$ \pi$ CC π C π CCAG	CTGGGGGAGC	CCTGCAAGCA	CCCGGGGTCC	1920
MACCONC TO CO.	P GAGCAACCTG	. CCAGCCCGTG	CCACTGGTTG	TTTTGTTATC	MCTCICIAGG	1980
	, աստարարը արարարար	- $TCTCTCACTC$	GTTCATTCAT	CCAGGCATTC	WILLGWCWWII	2040
mammeacha)	¬ സസ አጥ አጥርጥርር	CAGACACCAG	AGACAAAATG	GTGAGCAAAG	CAGICACIGC	2100
COMMODERATION OF THE PROPERTY	* MCCXCCMCXC	• ልርጣጥጥርጥርልጥ	GGAAGACGTG	CAGAAGAAAA	TTAATAGCCA	2160
COCA A CITTA I	\ አሮሮሮ <u>ል</u> ሮሞርሮባ	' GAAAGAAAGG	AAATAAACAC	CATCTTGAAG	AATTGTGCGC	2220
A CO A MCCCCMI	ኮ አአሮአአርርርር	. CCTCCCTAGC	GCCCCCTGCT	GCCTCCATCG	TGCCCGGAGG	2280
00000033000	~	• AACCCTCCTC	CTCCATCAGT	CACAGCGCIG	CWGCIGGCCI	2340
A C C MC C C C M TV	- CCCTCAATCC	· TOOTGOTGO	TCTGAGCTGG	AGACTCCTTG	GCTCCAGGCT	2400
CC3 C3 3 3 CC	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	: GAAACTAGTC	TAACGGAGAA	TCTGGAGGGG	ACAGIGITIC	2460
OMONONCOC	እ <u>እ</u> እርርርርርርር	' CACGTCCAGG	AGAATTCCAG	GAGGTGGG	CIGCAGGGAG	2520
TCCCCACCC	T CCCCCTGAG	т ссстсстсаа	AGGCAGGAAG	GTGAAAAGGG	CAAGGCIGAA	2580
COMCOCCAC	3 MCMMC3CMC5	r ጥርጥጥር እ ርርርር	CCTGGGACTT	TTCCGTTGC1	TUCTUTUMGC	2640
	m mmmcmcmccc	P TOGAGGAGAA	GAAGTCTATT	TCATGAAGG	ATGCAGTTC	2700
AUD D D CALC	G CTGTTAAAA'	P TOCAGGGTGT	r GCATGGGTTT	TCCTTCACG	MGGCCTITAL	2760
WILL TANK OF CO	A TATAGGAAG	ጉ ርኔርርጥርኔጥጥ ባ	' CCTAGGCCG1	P TAATTCACG	AAGAAGIGAC	2820
	T TCTTTCATG	T CTTCTGGGC	ACTACTCAGO	CCTGTGGTG	3 ACTTGGCTTA	2880
mcca a ca co	C TCGAAAACC	TO TOGALATORGO	AGACTCGGTT	r TTCTTTCTG(TTCTGCCATT	2940
CCMMCCCMC	T CCCACCCTC	G GCAAGTGTCT	r CTCCTTCCC	r GGGCCATAG'	r CTTCTCTGCT	3000
AUD V V GOC 1G	C TTGCAGCTC	r cgrgrrcrg	GAACACTTC	CTGTGATTC	r CTGTGAGGGG	3060
WINWOVCC						

			GAGCAGCTGA	GCCACAGGGG	AGGTGGAGGG	3120
	AGGGGAAGGA		CCATCAGTCC	TCACTGATCA	CGTCAGACTC	3180
GGACAGGAAG	• • • • • • • • • • • • • • • • • • • •		GCTCAATGAA	CCCAACAGCC	ACATTTTCCT	3240
CAGGACCGAG	AGCCACAATG	CIICIO	TAACCAAAAA	GAATGCAGAG	ACTAACTGGT	3300
TCCCTAAGCA	TAGACAATGG	CAAAAACTGG		GTGGAAAATG	CCAGAGATTG	3360
GGTAGCTTTT	GCCTGGCATT		GCAGCTCAGC	AGTGACTGCT	GACAGCACGG	3420
TTAAACTTTT	CACCCTGACC	AGCACCCCAC AGGAGAAGAA		ATAGTGTATG	AGCAAGAAAG	3480
AGTGACCTGC	AGCGCAGGGG		CCACAGGGAT	TATAGTCCAC	GTGATCCTGG	3540
ACAGATTCAT	TCAAGGGCAG	TGGGAATTGA	GAAAAAATCA	GTTCAAGGGA	AGTCGGGAGA	3600
GTTCTAGGAG	GCAGGGCTAT	ATTGTGGGGG	CAAGCTGAGT	AATTCTGAGC	AAGTCACAAG	3660
CCTGATTTCT	AATACTATAT	TTTTCCTTTA	TTCTCCTTAT	TAGGAACTCT	TTTTCTCTGT	3720
GTAGTAACTG	AGGCTGTAAG	ATTACTTAGT	CTTTTAACAG	GAAGAAAACA	TTCCTAAGAG	3780
GGAGTTAGCA	GCACAAGGGC	AATCCCGTTT	GCTGACTATA	TGATTGGTTT	TTTGAAAAAT	3840
TAAAGCCAAA	CAGATTCAAG	CCTAGGTCTT	AAAATGAGAC	TAGTACCCTT	TGGTCAGCTG	3900
CATTTCAGCG		TCTGATTCAG	•		CCAATCAAAA	3960
TAAACAAACA	CCCAGTTGTA	AATGTCTCAA	CTCCACATCT		CTGCCAGGGC	4020
AGAATAGAAT	CTTTAGAGCA	AACTGTGTTT	ACACTGTTGT		AACATAAAGT	4080
AGTTTGGAAA	TATTTACTTC	ACAAGTATTG	TTAAAGTTAC		TTTGGTATAT	4140
TGCTCAAAGG	CAATCATTAT	TTCAAGTGGC			TGTAAAGCAG	4200
TTATTGGCTA		CTTTTTGTTT				4260
GGATTATTAA	CCTACAGTCC					4320
TGTTTTTACC	ACCTTCTAAC	TAAATTTAAC			·	4380
CTCAAAGTGG	TAATAACAGT					4440
TTATACTATA				• •		4500
CTTTGAAATT	AGACCTCCTG					4560
ATTTTGATAT	TTTGATAATC					4620
ATATATTTG	AAACATCTTT					4680
CATGCACAC	A CACAGAGTAA					4740
TGCAAGACT	AAATTAGAAA		_			4800
AGGGGGGAA	A TCTGCCGCTT				GGACTTGTTT	4860
CTTGTGTTC'	r GGCTGGCTG7				GTTTGCAGAG	4920
GGATCTCCA	G TTCCTAGCAT					4980
TGAATGGAA	A TATAAACTA					5040
TGTAAGTGT				• • • • • • • • • • • • • • • • • • • •	A AAAGAAACTC	5100
TAGGAACTA			• •		A AGGCAACCCC	5160
CAAACAGAC	T TCTGGAAGG				A TGAAGGGCTG	5220
CCTGTGCAC	A GCCCCACCC	A GCCTCACGT				
GCTCCCCAG	T ATATATAAA	C CTCTCTGGA	G CTCGGGCAT	G AGCCAGCAA		

(2) INFORMATION FOR SEQ ID NO:35:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 19 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:

AACTATTATT GGGGTATGG

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(2) INFORMATION FOR SEQ ID NO:36:

- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 19 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single

 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

TTGGTGAGGC TTCCTCTGG

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(2) INFORMATION FOR SEQ ID NO:37:

- (i) SEQUENCE CHARACTERISTICS:

 - (A) LENGTH:
 (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:

AACTAT TATTGGGGTA TGGGTGCATA AATTGGGATG TTCTTTTTAA AAAGAAACTC 5100 CAAACAGACT TCTGGAAGGT TATTTTCTAA GAATCTTGCT GGCAGCGTGA AGGCAACCCC 5160 CCTGTGCACA GCCCCACCCA GCCTCACGTG GCCACCTCTG TCTTCCCCCA TGAAGGGCTG 5220 GCTCCCCAGT ATATATAAAC CTCTCTGGAG CTCGGGCATG AGCCAGCAAG GCCACCCATC 5280 CAGGCACCTC TCAGCACAGC AGAGCTTTCC AGAGGAAGCC TCACCAA

(2) INFORMATION FOR SEQ ID NO:38:

- (i) SEQUENCE CHARACTERISTICS:

 - (A) LENGTH:
 (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:

AACTAT TATTGGGGTA TGGGTGCATA AATTGGGATG TTCTTTTTAA AAAGAAACTC 5100 CAAACAGACT TCTGGAAGGT TATTTTCTAA GAATCTTGCT GGCAGCGTGA AGGCAACCCC 5160 CCTGTGCACA GCCCCACCCA GCCTCACGTG GCCACCTCTG TCTTCCCCCA TGAAGGGCTG 5220 GCTCCCCAGT ATATATAAAC CTCTCTGGAG CTCGGGCATG AGCCAGCAAG G

INTERNATIONAL SEARCH REPORT

ational Application No. PCT/US 00/00559

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C12Q1/68 A61F A61P27/06 C12N5/10 C12N15/85 A61K31/7088 A61K48/00 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) C12Q A61P C12N Á61K IPC 7 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. Category NGUYEN TH D ET AL: "Gene structure and 1-111 X properties of TIGR, an olfactomedin-related glycoprotein cloned from glucocorticoid-induced trabecular meshwork cells" JOURNAL OF BIOLOGICAL CHEMISTRY, US, AMERICAN SOCIETY OF BIOLOGICAL CHEMISTS, BALTIMORE, MD, vol. 273, no. 11 13 March 1998 (1998-03-13), pages 6341-6350-6350, XP002132916 ISSN: 0021-9258 the whole document 1-111 WO 98 44107 A (UNIVERISITY OF CALIFORNIA X ;HUANG WEIDONG (US); NGUYEN THAI D (US);) 8 October 1998 (1998-10-08) the whole document Further documents are listed in the continuation of box C. Patent family members are listed in annex. * Special categories of cited documents : T* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the *A* document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. *O* document referring to an oral disclosure, use, exhibition or other means *P* document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 25 May 2000 31/05/2000 Name and mailing address of the ISA Authorized officer Ruropean Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016

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Müller, F

INTERNATIONAL SEARCH REPORT

ational Application No PCT/US 00/00559

		PC1703 00,	00333
C.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
X	WO 98 32850 A (UNIV CALIFORNIA) 30 July 1998 (1998-07-30) the whole document		1-111
X	WO 96 14411 A (UNIV CALIFORNIA) 17 May 1996 (1996-05-17) see whole doc. esp. claims		1-111
X	KUBOTA R ET AL: "Genomic organisation of the human myocilin gene (MYOC) responsible for primary open angle glaucoma" BIOCHEMICAL AND BIOPHYSICAL RESEARCH COMMUNICATIONS, US, ACADEMIC PRESS INC. ORLANDO, FL, vol. 242, no. 242, 1998, pages 396-400-400, XP002120488 ISSN: 0006-291X se whole doc. esp. fig. 2		70
Р,Х	WO 99 16898 A (ANCTIL JEAN LOUIS ;COTE GILLES (CA); MORISSETTE JEAN (CA); RAYMOND) 8 April 1999 (1999-04-08) the whole document		1-111

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